

(19)



Europäisches Patentamt  
European Patent Office  
Office européen des brevets

(11) Publication number:

0 297 723  
A2

(12)

# EUROPEAN PATENT APPLICATION

(21) Application number: 88304932.2

(51) Int. Cl. 4: A61M 1/10

(22) Date of filing: 31.05.88

(30) Priority: 29.05.87 US 56401  
13.11.87 US 120591  
25.05.88 US 198441

(43) Date of publication of application:  
04.01.89 Bulletin 89/01

(84) Designated Contracting States:  
DE FR GB IT SE

(71) Applicant: RETROPERFUSION SYSTEMS, INC.  
3178 Pullman Avenue  
Costa Mesa California 92626(US)

(72) Inventor: Lundquist, Ingemar H.  
17 Miles Drive at Dunes Road  
Pebble Beach California 93953(US)  
Inventor: Tarczy-Hornoch, Zoltan  
7106 Marlborough Terrace  
Berkeley California 94705(US)  
Inventor: Kardos, Thomas J.  
1570 Via Capri, No. 10  
Laguna Beach California 92651(US)

(74) Representative: Cross, Rupert Edward Blount  
et al  
BOULT, WADE & TENNANT 27 Fumival Street  
London EC4A 1PQ(GB)

(54) Retroperfusion and retroinfusion control apparatus, system and method.

(57) Retroperfusion control apparatus (11) for supplying arterial blood of a patient to the venous side of the patient's heart including a pump (81) having an inlet (104) and an outlet (114) and a piston (88) movable through a pump stroke for moving a liquid from the inlet (104) to the outlet (114) of the pump (81). A stepper motor (47) is provided which drives the piston (81). Electronic circuitry is provided for driving the stepper motor (47) and senses the presence of an R wave in a patient to operate the stepper motor (47) in response to the sensed R wave.

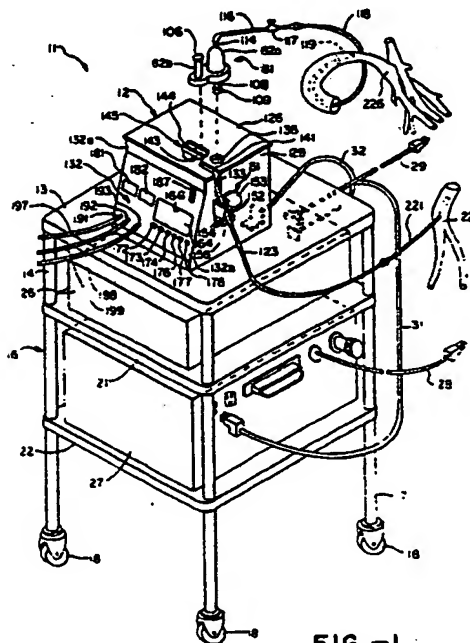


FIG. -1

EP 0 297 723 A2

## RETROPERFUSION AND RETROINFUSION CONTROL APPARATUS, SYSTEM AND METHOD

This invention relates to a retroperfusion, retroinfusion control apparatus, system and method.

This application is a continuation-in-part of application Serial No. 120,591, filed on November 13, 1987 which is a continuation-in-part of application Serial No. 056,401, filed on May 29, 1987.

Attempts have heretofore been made to perform synchronized diastolic coronary venous retroperfusion. Results are published in the August 1985 issue of the Journal of the American College of Cardiology, Vol. 6, No. 2, pages 328-335 and in Circulation, August 1986, Vol. 74, No. 2, pages 381-388. Both of these articles describe work which was done in connection with a synchronous retroperfusion system (USCI Model ECI). Such a system consists of a Hewlett Packard 78348A monitor for display of cardiac rhythm, arterial pressure and the pump signal. The monitor is a two-channel unit which is capable of monitoring and displaying the electrocardiogram and pressure or pump timing. Information from the monitor is fed and processed by the pump controller which operates a piston driven pump to maintain pump flow and pump timing through feedback circuits that compensate for variations in the patient's heart rhythm and rate. The piston driven pump is of the disposable type and is connected through tubing between the arterial blood supply and an auto-inflatable retroperfusion balloon catheter which is positioned in the great cardiac vein via the coronary sinus. As arterial blood is delivered through the coronary sinus catheter during diastole, this arterial blood inflates a balloon at the tip of the catheter. Inflation of the balloon seals the coronary sinus preventing leakage of arterial blood and permits a more effective retrograde delivery of arterial blood into the myocardium. On termination of retrograde catheter perfusion at or near end-diastole, the reverse stroke of the pump creates a back flow into and through the catheter which attempts to deflate the balloon. The amount of balloon deflation is heart rate and flow rate dependent. This allows retrograde coronary sinus drainage of venous blood from the myocardium into the right atrium during systole. Even though such work has been carried out in connection with retroperfusion, there is a need for a new and improved apparatus and system for carrying out such retroperfusion and an improved method for accomplishing the same. Essentially a retroperfusion control apparatus, system and method is disclosed in application Serial No. 056,401 filed on May 29, 1987. In addition, it has been found that it is desirable to provide an apparatus, system and method which can be utilized for retroinfusion and

which does not require the use of autoinflatable balloons. There is therefore a need for an improved apparatus and system which can be utilized for carrying out retroinfusion as well as retroperfusion.

In general, it is an object of the present invention to provide a retroperfusion and retroinfusion control apparatus, system and method which makes it possible to accomplish retroperfusion and retroinfusion in humans more proficiently.

Another object of the invention is to provide a retroperfusion and retroinfusion apparatus, system and method of the above character in which adjustable delivery rate by stroke length and timing of pumping is utilized which is synchronized to the heart's R-wave signals.

Another object of the invention is to provide a retroperfusion and retroinfusion control apparatus, system and method of the above character in which the pump cycle is always terminated at or before the beginning of a new R-wave.

Another object of the invention is to provide a retroperfusion and retroinfusion control apparatus, system and method which utilizes a microcomputer for monitoring the R-waves for initiating and terminating the pump cycle.

Another object of the invention is to provide a retroperfusion and retroinfusion control apparatus, system and method of the above character which can accommodate irregular heartbeats in the patient.

Another object of the invention is to provide a retroperfusion and retroinfusion control apparatus, system and method of the above character in which a computer controlled stepping motor is utilized for providing an adjustable delivery rate such as by adjustable pump stroke and adjustable stepping rate.

Another object of the invention is to provide a retroperfusion and retroinfusion control apparatus, system and method in which it is possible to specify delivery volume/time and/or delivery pressures.

Another object of the invention is to provide a retroperfusion and retroinfusion control apparatus, system and method of the above character in which the pumping can be controlled with great precision.

Another object of the invention is to provide a retroperfusion and retroinfusion control apparatus, system and method which utilizes a powered downstroke or rearward stroke as well as powered upstroke or forward stroke.

Another object of the invention is to provide a retroperfusion and retroinfusion control apparatus, system and method in which an active precisely controlled vacuum stroke of the pump motion is

provided to accentuate deflation of the blood inflated balloon.

Another object of the invention is to provide a retroperfusion and retroinfusion control apparatus, system and method in which the ECG signal from the patient is differentiated to find the maximum positive slope of the ECG waveform to provide an independent signal that an R wave is occurring or arterial pressure is differentiated to find a maximum negative slope to provide an independent signal that diastole is beginning.

Another object of the invention is to provide a retroperfusion and retroinfusion control apparatus, system and method in which it is possible to more precisely ascertain when the R wave is occurring.

Another object of the invention is to provide a retroperfusion and retroinfusion control apparatus, system and method in which a reviewing action is utilized in the pump to facilitate deflation of the blood inflated balloon.

Another object of the invention is to provide a retroperfusion and retroinfusion control apparatus, system and method which utilizes a pinch-off valve for shutting off blood flow during non-pumping modes.

Another object of the invention is to provide a retroperfusion and retroinfusion control apparatus, system and method of the above character which utilizes a blood level sensor to detect blood supply problems such as an air leak or an occluded supply catheter or tubing.

Another object of the invention is to provide a retroperfusion and retroinfusion control apparatus, system and method of the above character which incorporates numerous safety features.

Another object of the invention is to provide a retroperfusion and retroinfusion control apparatus, system and method of the above character in which it is possible to deliver greater quantity of oxygenated blood to the area at risk even though the patient may have complex arrhythmias.

Another object of the invention is to provide a retroperfusion and retroinfusion control apparatus, system and method of the above character in which the timing can be varied independently of blood flow to allow optimal perfusion of the myocardium.

Another object of the invention is to provide a retroperfusion and retroinfusion control apparatus system and method of the above character in which the blood of the patient is utilized for autoinflating the balloon or alternatively, a fluid is introduced exterior of the patient's body for inflating the balloon.

Another object of the invention is to provide a retroperfusion and retroinfusion control apparatus system and method of the above character in which gas is utilized for inflation of the balloon.

Another object of the invention is to provide a retroperfusion and retroinfusion control apparatus, system and method in which a central control console is utilized having an electroluminescent flat panel display.

Another object of the invention is to provide a retroperfusion and retroinfusion control apparatus, system and method of the above character in which the sinus pressure is utilized to automatically control the pumping of blood and/or the balloon inflation.

Another object of the invention is to provide a retroperfusion and retroinfusion control apparatus, system and method of the above character in which an electronically isolated ECG input capability is provided.

Another object of the invention is to provide a retroperfusion and retroinfusion control apparatus, system and method of the above character in which electronics is provided which makes it possible to connect ECG electrode pads directly to the patient.

Another object of the invention is to provide a retroperfusion and retroinfusion control apparatus, system and method which makes it possible to maintain inflation of the catheter balloon for more than one cardiac cycle.

Another object of the invention is to provide a retroperfusion and retroinfusion control apparatus, system and method of the above character which can be used in cerebral veins as well as in cardiac veins.

Another object of the invention is to provide a retroperfusion and retroinfusion control apparatus, system and method of the above character in which non-synchronous gas inflation for more than one cardiac cycle can be accomplished.

Another object of the invention is to provide a retroperfusion and retroinfusion control apparatus, system and method of the above character which utilizes a cassette which has an intermediate captive driving solution.

Additional objects and features of the invention will appear from the following description in which the preferred embodiment is set forth in detail in conjunction with the accompanying drawings.

Figure 1 is an isometric view of a retroperfusion control apparatus and system incorporating the present invention and showing the disposable pump about to be inserted into the same.

Figure 2 is a front elevational view of the controller shown in Figure 1.

Figure 3 is a side elevational view of the apparatus shown in Figure 1.

Figure 4 is a front elevational view of a disposable pump incorporating the present invention.

Figure 5 is a side elevational view in cross section of the pump shown in Figure 4.

Figure 6 is a block diagram of the electronics used on the system incorporating the present invention.

Figures 7A, 7B, 7C and 7D are strip chart recordings showing test results of the apparatus of the present invention at different heart beat rates.

Figure 8 is an isometric view similar to Figure 1 but modified to incorporate additional improvements and in particular to incorporate the capability of performing retroinfusion and gas inflation of the balloon.

Figure 9 is a side elevational view of the apparatus shown in Figure 8.

Figure 10 is a schematic illustration of the portion of the apparatus utilized for inflating the balloon with a gas.

Figure 11 is a chart showing a timing diagram for the gas inflation apparatus shown in Figure 10.

Figure 12 is a block diagram of the electronic circuitry utilized for controlling the gas inflation apparatus shown in Figure 10.

Figure 13 is a chart showing the timing diagram for the circuitry shown in Figure 12.

Figure 14 is a schematic diagram of an alternative air inflation mechanism for the balloon.

Figure 15 is a block diagram of the electronic circuitry for controlling the apparatus shown in Figures 8 and 9.

Figure 16 is an isometric view of a control apparatus and system similar to that shown in Figures 1 and 8 but modified to incorporating the use of a bladder pump cassette and a second gas balloon catheter for cerebral retroperfusion.

Figure 17 is a front elevational view of the controller shown in Figure 16.

Figure 18 is a front elevational view partially in cross-section of a cassette for use in the control apparatus shown in Figure 16.

Figure 19 is a side elevational view partially in cross section of the cassette shown in Figure 18.

Figure 20 is a cross sectional view taken along the line 20-20 of Figure 19.

In general, the retroperfusion and retroinfusion control apparatus is comprised of a positive displacement pump having an inlet and an outlet, piston-like means for moving liquid from the inlet to the outlet, a stepper motor, means coupling the stepper motor to the piston-like means for causing operation of the piston-like means, electronic circuitry for driving the stepper motor, the electronic circuitry including means for sensing the R wave of a patient for operating the stepper motor, and for displaying the electrocardiogram, R wave, the pump stroke timing in relation to the above.

More in particular, the retroperfusion and

retroinfusion control apparatus and system 11 consists of a pump console 12 which is supported on the top level of 13 of an equipment dolly or stand 14. The equipment dolly or stand 14 is provided with a rectangular framework 16 which has four depending legs 17 having casters 18 mounted on the bottom extremities of the same. The equipment dolly or stand 14 is provided with an intermediate level shelf 21 and a lower level shelf 22 which are carried by the legs 17.

A power supply 26 for the pump console 12 is mounted upon the intermediate shelf 21. A backup power supply 27 of the battery type is mounted on the lower shelf 22. The backup power supply 27 is provided with a conventional electrical cord 28 which is adapted to be connected to a conventional source of ac, as for example, 110 volts 60 cycle ac. The power supply 26 is provided with a similar electrical cord 29 which also is adapted to be connected to either a conventional type ac outlet or the backup power supply 27. Another electrical cord or cable 31 is provided which interconnects the backup power supply 27 to the power supply 26. A cord 32 connects the power supply 26 to the pump console 12.

The pump console 12 is provided with an internal metal framework 36 which is divided into a pump drive compartment 37 a monitoring compartment 38 which is positioned below the pump drive compartment 37 and pc board compartments 39 and 41 which are mounted on the other side of the framework 36. The pump drive compartment 37 is provided with a support plate 46 which forms a part of the framework 36. A stepping motor 47 is mounted on the support plate 46 and is provided with an output shaft 48 that extends through the support plate 46. Another support plate 56 is provided which is mounted upon posts 57 carried by the support plate 46. A rack 61 is mounted for vertical reciprocation in a bracket or slide 62 which is mounted upon the support plate 56. A pinion 63 engages the rack 61 and is mounted upon a shaft 64 that is carried by a coupling 66 mounted upon the output shaft 48.

Yieldable spring means reduces the drive system compliance by preloading rack 61 in tension and consists of a spring 71 which has one end connected to a pin 72 which travels with the rack 61 and which has the other end connected to a pin 73 which is mounted upon the support plate 56. Motor position switch means 75 is provided for giving a timing signal when the rack 61 has reached its lowermost position and consists of infrared sensing means in the form of a light emitting diode 76 and a photosensor 77 carried by a bracket 78 mounted on the plate 46. A vane 79 mounted on the coupling 66 is adapted to pass between the diode 76 and the photosensor 77 to provide the

timing signal.

A disposable pump cassette 81 adapted to be utilized with the pump console 12 is shown in Figures 4 and 5 and consists of a pump body 82 which is provided with two portions 82a and 82b in which portion 82a is dome-shaped and forms a pump chamber 83 and portion 82b is semi-cylindrical and forms a bubble chamber 84. A pump base 86 is secured to the lower extremity of the pump body 82 by suitable means such as ultrasonic welding. The pump base 86 is provided with a cylindrical depending open ended portion 87 which opens into the chamber 83. A piston 88 is mounted for reciprocatory movement within the cylindrical portion 87 and extends upwardly into the chamber 83. A boot 89 of a suitable material such as a silicon rubber is positioned over the piston 88 and has its lower outer margin secured between the pump base 86 and a boot retainer 91.

An O-ring is disposed below the boot 89 and is in sealing engagement with the cylindrical surface of the piston 88 as shown particularly in Figure 5. A seal member 93 of a suitable material such as silicone rubber is disposed in the pump chamber 83 and has its outer margin clamped between the pump body 82 and a retainer 94. An offset flapper valve member 95 as shown is provided with a tapered construction so that its thickness decreases progressively towards the inner margin of the same. The valve member 95 overlies a flow passage 96 which is in communication with a flow passage 97 that extends into a semi-annular flow passage 98 opening into the bubble chamber 84.

The boot retainer 91 is provided with a cylindrical upstanding portion 101 which opens into the interior of the bubble chamber 84. The cylindrical portion 101 is provided with a flow passage 102 which extends downwardly through the same and through the pump base 86 to inlet connection 104. A resealable membrane cap 106 formed of a suitable material such as rubber is mounted on top of the portion 82b of the pump body 82 and encloses the chamber 84. A plunger or lower piston rod 108 is provided which is secured to the piston 88 and depends downwardly therefrom. The piston rod 108 is provided with a head 109 which is adapted to seat within a recess 111 provided in the upper extremity of the rack 81. A retaining cap 112 having an opening 113 through which the piston rod 108 extends is secured to the lower extremity of the cylindrical portion 87. A protrusion 114 is provided on the upper extremity of the pump body 82 and is provided with a flow passage 116 therein which opens into the pump chamber 83.

In order that the pump cassette be disposable, it is desirable that the pump cassette be formed of inexpensive materials. With the exception of the boot 89, the seal and valve member 93, the o-ring

92 and the cap 106, all of the remaining parts can be formed of suitable materials, such as plastic. The use of clear plastic makes it possible to see into the pump chamber 83 and into the bubble chamber 84.

The protrusion or outlet 114 is sized so that a tubing 116 can be bonded to it. The tubing 116 is provided with a clamp 117 and is adapted to be connected to an auto-inflatable retroperfusion balloon catheter 118 of a conventional type which is introduced into the venous side of the heart during retroperfusion operations as hereinafter described. The catheter 118 is provided with an inflatable balloon 119. The protrusion or inlet 104 is sized so that the tubing 123 which extends downwardly through a compartment 124 can be bonded to it. The tubing 123 is adapted to be connected to an arterial blood supply as hereinafter described. The tubing 123 is sufficiently compliant so that the lumen extending therethrough may be occluded by the jaws of the pinch valve 154.

The pump console 12 is provided with a top wall 126 which forms a part of the framework 36. It is also provided with a bottom wall 127, a rear wall 128, side walls 129 and 131 and a front wall 132. The front wall 132 is provided with an upper vertical portion 132a and lower inclined portion 132b. The compartment 124 opens to the exterior through a slot 133 in the side wall 129. The top wall 126 is provided with an opening 136 through which the lower extremity of the disposable pump cassette 81 can extend and in particular so that the retaining cap 112 can engage lips 137 which extend into the opening 136 at the time that the head 109 is slid into the recess 111. Yieldable means is provided for retaining the disposable pump cassette 81 in a predetermined position on the top wall 126 and includes a pair of spring urged detents 138 which are carried by the top wall 126 adjacent the opening 136 and which are adapted to engage dimples 139 provided on the bottom side of the pump base 86.

The top wall 126 is provided with another slot 141 which is adapted to receive the protrusion 104 and the tubing 123 carried thereby. As the pump cassette 81 is moved into place, the portion 82b of the pump body 82 forming the bubble chamber 84 is moved in between a pair of spaced apart members 143 which are mounted on a post 144 provided on the top plate or wall 126.

Infrared sensing means 145 is provided for sensing the level of liquid in the bubble chamber 84 and consists of a pair of light emitting diodes 146 and photodiodes 147 which are carried by the members 143. One light emitting diode 146 is provided on each of the members 143 and one photodiode 147 is provided on each of the members with the photodiodes facing the light emitting

diodes. The photodiodes are provided to sense when the level of the blood within the chamber 84 drops below a predetermined level to stop the pumping action as hereinafter described.

A pinch off valve mechanism 151 for cutting off arterial blood flow through the tubing 123 is provided on the side wall 129 of the pump console 12 and consists of a bracket 152 which is secured to the side wall 129. An electrical solenoid 153 is mounted on the bracket and is adapted to operate a clamping mechanism 154 provided on the bracket 152. The clamping mechanism 154 is of a conventional type and is spring loaded in an open position and is provided with a slot 156 therein through which the tube 123 can extend. When the solenoid 153 is energized, the slot 156 is closed to pinch off flow through the tubing 123. As soon as the solenoid 153 is de-energized, arterial blood flow resumes through the tubing 123.

Cassette position sensing means 158 is provided for ascertaining when the pump cassette 81 is properly positioned on the pump console 12. This sensing means 158 consists of a microswitch 159 mounted on the top plate 126 (see Figure 2). It is provided with an operating arm 160 which engages the retaining cap 112 of the cassette 81.

A cathode ray tube 162 is mounted in the monitoring compartment 38 and is seated upon an inclined plane 163 mounted within the framework 36. The cathode ray tube 162 is provided with a screen 164 which is visible through an opening 166 provided in the front panel 132. A keyboard 171 is provided below the screen 164 and is provided with 6 push buttons 172, 173, 174, 176, 177 and 178.

Push button 172 is a spare. Push button 173 which carries an up-arrow function and push button 174 which carries a down-arrow function are utilized for increasing and decreasing the number of the various numerical parameters which are utilized for controlling operation of the pump cassette 81 as hereinafter described. The fourth key 176 is a start/stop key and controls the starting and stopping of the pump. The key 177 is the menu key and can be utilized for toggling between the ECG display screen and the numeric menu screen. The sixth key 178 is the line key. In addition to the main CRT screen 164 that can be viewed from the front panel 132, two three digit light emitting diode displays 181 and 182 are provided. Display 182 is for the flow rate setting for the apparatus and display 181 is for displaying the heart rate which is being sensed from the electrocardiogram.

Another display on the front panel shows the travel of the rack 61 and is comprised of a light emitting diode 184 supported on a bracket 186 mounted on the rack 61 so that it travels vertically with the rack. The travel of the light emitting diode

184 is visible through a slot 187 provided in the front panel 132.

Three connectors 191, 192 and 193 are provided in the lower left-hand side of the front panel 132. The connector 191 is used for receiving the electrocardiogram signal output which typically is a one volt peak-to-peak signal coming from an external ECG monitor. The connector 192 is utilized for providing the pump stroke signal which can be utilized in a strip chart recorder for externally recording the operation of the apparatus. The third connector 193 provides a piston position signal which can be utilized in a strip chart recorder for externally recording piston 108 position. This piston position signal is supplied by a linear potentiometer 194 having a movable armature 195 connected to piston 108 by an arm. Alternatively, the third connector 193 can be utilized as an input for measuring pressure, for example, pressures in the coronary sinus.

Conducting cords or cables 197, 198 and 199 (see Fig. 1) are connected to the connectors 191, 192 and 193 respectively. As shown in the block diagram in Figure 6 showing the electronics for the apparatus and system of the present invention, the cord or cable 194 is connected to an external patient ECG monitor 201. The patient ECG monitor 201 is of a conventional type and is provided with conventional leads 202 which are connected to electrodes 203 which are secured to the patient's body in appropriate locations in a conventional manner.

The pump console 12 contains numerous printed circuit boards which contain the circuitry which is shown in Figure 6. Thus there is provided a microprocessor and video controller board 206 which incorporates an analog-to-digital converter 207. The output from the patient ECG monitor 201 is supplied through a pre amp 208 which supplies its output to the analog-to-digital converter 207 and also to an ECG amplifier and processing circuitry 209. The circuitry 209 provides an "R" trigger signal to the microprocessor video controller 206. The DC power supply 26 is connected to various components of the circuitry as shown in Figure 6 as well as to the microprocessor and video controller 206. A motor interface and amplifier board 211 is provided which is used for controlling the stepper motor 47. Various other components of the pump console 12 are interconnected to the microprocessor video controller 206 as shown. These components include the keyboard 171 as well as the heart rate display 181 and the flow display 182. The motor position detector or switch means 75 associated with the stepper motor 47 is connected to the microprocessor and video controller 206 as well as to the motor interface and amplifier board 211. The infrared blood level detector 145 is con-

nected through a preamplifier circuit 212 through an A/D converter circuit 213 to the microprocessor 20. The cassette position sensing switch 158, the arterial pinch off valve assembly 151 and the CRT monitor 162 are also connected to the microprocessor and video controller 206. The microprocessor 206 supplies an output to the connector 192 which can be connected to a strip chart recorder. If desired, as shown in Figure 6, an additional external CRT monitor can be utilized.

Operation and use of the retroperfusion, apparatus and system in performing the method of the present invention may now be briefly described as follows. Let it be assumed that a patient has been identified in which it is desired to utilize a retroperfusion procedure. The equipment dolly or stand 14 is brought to the patient or conversely, the patient is brought to the equipment dolly where it is located. The cord 28 or 31 is connected into an appropriate power outlet in the hospital. The operator then observes the position of the rack 61 by noting the location of the small light emitting diode 184 which is viewable through the slot 187 provided on the front panel 132. This diode 184 should be at the bottom or home position. If it is not in the home position, the piston 108 should be shifted in the cassette 81 so that it is in an appropriate position so that it can be inserted into the pump console 12. The pump cassette 81 can then be positioned so that the retaining cap 112 enters the opening 136 so that the head 109 carried by the piston plunger 108 can enter the slot or recess 111 provided at the top of the rack 61. At the same time this is occurring, the bubble chamber is introduced into the infrared level detector 145. The on/off switch (not shown) for the power supply 26 can be operated to supply power to the control pump console 12. The screen 164 of the TV monitor should then be viewed to see whether or not the top trace 216 on the screen is a straight line. This trace 216 represents the pump stroke when the pump is being operated.

The cable 194 is then connected to the patient ECG monitor 201 and the ECG electrodes 203 are attached to the patient. The R wave signal on the video monitor 162 is then observed by observing the lower trace 217 on the screen 164 of the video monitor. The rate of rise of the R wave should always be greater than that of the T wave. Appropriate selection of ECG leads can be made on the patient to adjust the quality of the ECG signal. The cable 196 can then be connected to the strip chart recorder for recording the pump stroke cycle trace.

With an empty pump cassette in place, the start button 176 on the keyboard 171 can be pressed which will cause operation of the stepper motor 47 to cause operation of the rack 61 to operate the plunger 108 of the cassette. Unless

blood level sensor 212 senses an appropriate blood level, only one pump stroke is taken and the pump stops, indicating an alarm condition.

The pump console 12 may now be set to the desired parameters. For example, an initial flow rate of 100 milliliters per minute per EKG can be set by pressing the menu button 177 to cause the menu screen to be displayed. The up and down keys 173 and 174 can then be utilized to select the correct value for flow. As soon as these values have been set the menu button 177 can be again pressed to enter the new value and execute the flow adjustment. The screen will again display the ECG signal and the pump stroke trace.

Utilizing conventional aseptic techniques, a conventional supply catheter 221 can be placed in the femoral artery 222 of the patient and connected to the tubing 123 to obtain a supply of arterial blood. The inflatable retroperfusion balloon catheter 118 is placed in the great cardiac vein 226 approximately 2 to 3 centimeters proximal to the anterior interventricular vein via the coronary sinus. Proper placement of the catheter 118 is confirmed under fluoroscopy by observing the free flow of a radiopaque solution around the catheter in the atrium during systole. A disposable pressure transducer (not shown) can be secured to the proximal end of the retroperfusion catheter's pressure lumen if it is so equipped, and can be connected to the cabling 197 which is connected to the pressure connector 193 provided on the front panel 132.

A sterile pump cassette 81 is placed on the pump console 12. The apparatus is then primed with a sterile heparinized saline solution. Any air bubbles which appear in the apparatus are removed by using a syringe to penetrate the cap 106 and withdrawing air from the bubble chamber 84. In order to ensure that all air is withdrawn from the apparatus, a syringe can also be utilized to draw blood back through the outlet tubing 117 and retroperfusion catheter into the injection site 117a to ensure that all air has been removed.

As soon as this has been accomplished, the pump can be placed in operation. As arterial blood is delivered through the coronary sinus catheter during diastole, the blood automatically inflates the balloon at the end of the auto-inflatable retroperfusion balloon catheter 122. This retards the efflux of blood from the regional coronary veins and permits effective retrograde delivery of arterial blood to the myocardium of the heart. Termination of the retrograde catheter perfusion at end of diastole, automatically deflates the balloon and permits prograde coronary sinus drainage of venous blood from the myocardium into the right atrium during systole.

More specifically, the operation of the apparatus and in particular, the circuitry which is shown in

Figure 6 may now be described. The one volt peak-to-peak signal which is supplied from the external ECG monitor 201 has been prefiltered to approximately 150 hertz high frequency rolloff. A signal is supplied to the preamplifier 208 which accomplishes additional filtering and also supplies a signal to the A/D converter 207 connected to the microprocessor 206. A signal is also supplied from the preamp 208 to the ECG amplifier and processing circuitry 209 which is utilized to ascertain location of the R wave peak. This is accomplished by taking the electronic derivative of the signal to find the maximum slope in the wave form. This information is supplied as a signal on the R trigger circuit to the microprocessor 206. The microprocessor 206 has the capability of correlating the pattern which is generated by the A/D converter 207 based upon the ECG monitor and compares it with the R trigger circuit signal being supplied by the ECG amplifier and processing circuitry 209 to ascertain whether or not there is an agreement that an R wave has been detected which can be utilized for triggering the operating of the stepper motor 47.

The output of the microprocessor video controller 206 feeds signals to the stepper motor 47 that are phased in relationship to the stepper motor to create approximately 500 steps per inch of operation of the motor in forward and then in reverse and causing a resultant travel of the piston 88 in the pump cassette 81 between the upper and lower limits of movement for the piston 88. These steps of the stepper motor, and the rate at which these steps are taken, are controlled by the microprocessor 206 utilizing a lookup table and are based upon the heart rate input that is sensed by the ECG monitor 201 and the delivery setting which has been inserted by the physician into the pump console 12 by operation of the menu key 177 and the up and down keys 173 and 174 as hereinbefore described. In this manner, the microprocessor 206 determines precisely the upstroke time and speed as well as the pause time and the downstroke time and speed. The microprocessor 206 is capable of assimilating arrhythmias and abnormal ECG events. Typically the microprocessor 206 initiates the pump cycle at approximately 45% and terminates at 95% of the R to R period if the heartbeat is steady. The desired steadiness can be defined, e.g. no more than 10% change during the past eight heartbeats. If the rate is changing more rapidly, the pump is started later depending upon the rate of change to avoid premature pumping and straining of the heart. The pump cycle is always terminated at the beginning of a new R wave. After a very irregular beat the pump skips a pump cycle until the heartbeat stabilizes, all under the control of an algorithm. As hereinbefore explained there are three feedbacks to the microprocessor from the

stepper motor 47, from the pump cassette 81, and from the infra-red blood level detector. The first is the bottom of stroke indication from the motor position detector 75. The second input to the microprocessor 206 is the cassette position switch 158 which informs the microprocessor whether or not the cassette is actually connected to the stepper motor rack 61. Thus, if the cassette is improperly placed, the microprocessor 206 will stop the pumping operation. Information is also supplied from the infrared blood level detector 145 to the microprocessor 206 and causes the microprocessor 206 to shut down the pumping action when air bubbles are sensed in the blood or when the blood level within the bubble chamber 84 falls below the level of either of the infrared sensors of the blood level detectors 145.

The microprocessor 206 has been programmed so that the pump stroke will start at 45 % of the R to R interval (the R period) and terminates at approximately 95% of the R to R interval. In programming this pumping operation, it has been found it is desirable to program the microprocessor 206 so that signals to the stepper motor 47 are entered twice as fast on the upstroke so as to leave approximately one half of the time allotted for the upstroke for a pause time after which the downstroke is commenced. The downstroke is accelerated in order to cause better deflation of the auto inflatable balloon 119 at the end of the catheter 118 and also to more quickly reduce the pressure from the coronary sinus caused by the pump stroke.

Such a flow pattern is shown in the traces shown in Figure 7. The traces show that pumping into the coronary sinus actually begins at the end of systole or in other words at the end of the arterial pressure wave. Utilizing a pause between the upstroke and downstroke, the slope of the rise of flow is much steeper and achieves peak flow much earlier in the R to R cycle which makes it possible to achieve peak pressure in the coronary sinus earlier in the R to R cycle. This helps to prevent or avoid the generation of overpressures or collisions between the pressure due to flow caused by the pump and the pressure wave due to arterial pressure of the next systole. Thus it can be seen that this helps to avoid possible hemorrhaging in the heart. A downward or receiving portion of the flow occurs prior to the occurrence of the next R wave, thus pressure is removed more rapidly from the coronary sinus.

With such use of a pause phase, the same amount of blood can be delivered between the R to R peak but is delivered more rapidly and more appropriately in the diastolic time window. In addition the pressure is removed more rapidly to prevent any possible collision from the systolic arterial pressure wave. By adding the pause at the end of

the upstroke, the pressure generated completely depletes and translates into maximum blood flow. The downstroke does not occur until after this pressure has been completely dissipated through the catheter and maximum flow has been expelled through the catheter. As the downstroke occurs, a greater negative pressure is created than would be the case without a pause. This greater negative pressure facilitates collapse of the balloon 119 of the auto-inflatable balloon catheter 118.

Thus it can be seen there are two effects from such a procedure. One is to deliver the pressure wave and the flow earlier in the R to R cycle and the other is to allow greater flow through the catheter by allowing the buildup of pressure to deplete itself and translate into greater blood flow with a subsequent improved collapse of the balloon.

In Figures 7A, 7B, 7C and 7D, there are shown four strip chart recordings which show the response of the pump 81 in response to heart rate variation. In achieving the data which is shown in Figures 7A, 7B, 7C and 7D, a retroperfusion apparatus and system of the present invention was utilized with the pinch valve assembly 151 being utilized for controlling the arterial blood flow. A patient ECG simulator was utilized to provide a normal rhythm. A conventional flow meter and a strip chart recorder were utilized. A saline solution was placed in a bag at 6 feet in elevation to simulate arterial pressure. A 7 French catheter with a 5 millimeter balloon was utilized. The catheter tip was introduced into a graduated cylinder so that the amount of saline solution which was pumped could be measured. In carrying out the tests, a calibration was performed for 0 to 100 milliliters per minute flow to calibrate the strip chart versus the graduated cylinder. In each of the flow settings of 20 to 120 milliliters per minute at 20 milliliter increments, the delivered mean flow was recorded on the strip chart recorder at heart rate settings of 60, 80, 120 and 150 beats per minute. Recordings were made at both slow and fast recording speeds with each setting as shown in Figures 7A, 7B, 7C and 7D. After the calibration had been completed, the flow was set at 80 milliliters per minute and the heart rate was varied through 40, 60, 80, 100, 120 and 150 beats per minute at a fast strip chart speed and then varied through 150, 120, 100, 80, 60 and 40 beats per minute at slow speed for compressed recording. The results of the tests are shown in the traces in 7A, 7B, 7C and 7D.

The trace 231 in Figure 7A, shows the electrocardiogram with the R peaks 232 as they are changed from a rate of 150 beats per minute to 120, 100, 80, 60, and 40 beats per minute. The phasic timing of the pump stroke is shown by the trace 236 in Figure 7B. The highest level 237 of the trace indicates the upstroke time. The next lower

level 238 indicates the duration of time that is occupied by the pause state where the pump piston is at its highest position and is held there for a predetermined period of time. The next level 239 indicates the time taken for the downstroke. The lowest level 241 indicates the pause or waiting time before the next trigger signal arrives to start the upstroke.

The trace 246 which is shown in Figure 7C shows the actual pump piston position with respect to time. The movement of the piston during the upstroke is indicated by the upwardly sloped portion 246a of the trace 246. The pause for the piston at the upper limit of its travel is indicated by the flat portion 246b and the downwardly inclined slope portion 246c indicates the downward stroke of the piston. The flat portion 246d represents the pause before the next upstroke of the piston is started.

The trace 251 which is shown in Figure 7D shows the time-averaged mean flow output from the pump through a calibrated flow meter and shows that a substantially constant output flow as, for example, 80 milliliters per minute for which the pump console 12 was set is achieved even though the beats per minute change radically. The trace in Figure 7C show how this was accomplished. As the beats per minute decreases, as for example, 150 for the initial pump stroke as shown by the level portion 246b, which is represented by the level 247 for 150 beats per minute. As the heart rate decreases, a larger volume of blood must be pumped with each stroke and therefore the stroke length is increased as represented by level 248 for 120 beats per minute, level 249 for 100 beats per minute, level 251 for 80 beats per minute, level 252 for 60 beats per minute and level 253 for 40 beats per minute. The trace 251 in Figure 7D shows that the output of the pump remains substantially constant through the entire operating range from 150 to 40 beats per minute. Thus it can be seen that the microprocessor 206 senses the change of rate of the heart beats and adjusts the upstroke time, the upstroke speed, the pause time and the downstroke time so that with the reduced number of strokes per minute increased volume is produced by the pump each time a stroke is made so that the resultant mean flow from the retroperfusion apparatus is substantially constant.

It has been found that once the system has been primed, there is no accumulation of air within the system because the system is sealed.

From the foregoing, it can be seen that there has been provided a retroperfusion apparatus system and method which has many advantages. The microprocessor controlled stepper motor drive provides a positive control over the pump stroke and provides a powered upstroke and a powered downstroke by forward and reverse motion of the step-

per motor. The precisely controlled powered down-stroke contributes to the balloon deflation. The microprocessor control which is utilized makes it possible to precisely detect the R waves by ascertaining the maximum positive slope within the ECG waveform and supplies a signal which is correlated with software in the microprocessor to ascertain whether in fact an R peak has occurred to therefore make possible a more positive and precise identification of the R wave. A direct coupling is provided between the stepper motor and the piston of the pump which direct coupling is obtained by the use of a rack and pinion.

Numerous safety features have been provided in the apparatus and system. In addition, the pinch-off valve in the arterial line clamps off the arterial line when the system is stopped or a fault alarm or condition occurs. This prevents flow through of the arterial blood. Thus, it can be seen that the pinch-off valve prevents passive flow through of arterial blood under arterial pressure through the system from the arterial side to the venous side. If the pinch-off valve were not present, it would be possible for such passive arterial blood flow to be as much as 30 milliliters per minute which could eventually fill the auto inflatable balloon and occlude the sinus for egress of blood which could have very deleterious effects on the patient.

Another embodiment of the retroperfusion and retroinfusion apparatus and system is shown in Figures 8-15 and is particularly adapted to perform retroinfusion as well as retroperfusion and which is adapted to be utilized with a gas inflated balloon which is inflated from a gas source external of the body, rather than being auto-inflated with the patient's blood.

As can be seen from the apparatus shown in Figures 8 and 9, major portions of the apparatus remain the same as shown in the previous embodiment of the invention. However, certain changes have been made. For example, the CRT display as represented by the cathode ray tube 162 has been removed and the space which was utilized by the same has been used to incorporate a power supply 261 which takes the place of the separate power supply 26 shown in Figure 1. This makes the pump console 12 self-contained except for the use of the backup power supply 27. Also in place of the CRT, an electroluminescent flat panel display 266 has been provided. The flat panel display 266 can be of any suitable type, such as one produced by Finlux Corporation of Finland, having by way of example 512 x 256 dot resolution. In addition, the six pad keyboard comprised of the switches 172, 173, 174, 176, 177 and 178 have been eliminated and a touch panel 268 has been provided which has been laminated directly over the display panel 267. This touch panel 268 is substantially transpar-

ent and can be operated by touching it with a finger or pen or some other object, in particular places on the touch panel to cause it to serve as a keyboard input. As also shown in Figure 9, a gas inflation mechanism or apparatus for balloon inflation as hereinafter described is incorporated in the space previously occupied by the CRT and is disposed behind the electroluminescent display panel 266 and in the front of the power supply 261.

In order to provide retroinfusion capability for the apparatus and system, a container such as a flexible bag or a bottle 276 is provided which can carry a diluent or a drug liquid within the container 276. The container or reservoir 276 can be suspended in a conventional manner such as by utilizing a strap 277 carried by the container 276 and secured to a hook 278 carried by a support member 279 mounted upon a stand 281. Suitable conventional means (not shown) can be utilized for controlling the flow of the liquid contained within the container 276 and for supplying the same into a tube 283 which can be connected in a suitable manner to the tube 221 to permit the drug or diluent to be supplied in the blood for the coronary sinus in an ECG synchronized or a non-synchronized manner. Thus the drug or diluent is introduced into the pumping circuit by connecting the pump inlet line previously attached to the arterial catheter to the reservoir 276 containing the drug or diluent retroinfusate. For the non-synchronized mode, the pump mechanism is operated at a high frequency (as for example, 300 cycles per minute) with each cycle providing a small stroke (such as less than 1 cc) such that the result of pulsatile flow approximates a continuous flow. Alternatively for low flows a slow stroke over a long time duration is provided while the pinch valve 153 is closed, with a fast filling stroke while the pinch valve 153 is open. These allow utilization of a pulsatile type pumping cassette for generating near continuous flow. In the synchronous approach, the flow is controlled in conjunction with the sensed sinus pressure and independently of the balloon inflation/deflation timing.

In the present embodiment, rather than using an auto-inflatable balloon, gas inflation means is provided for inflating and deflating the retroperfusion catheter balloon 119 in synchronism with each pump stroke or alternatively, in response to sensed sinus pressure. This gas inflation mechanism 271 is shown in Figure 9 and schematically in Figure 10. As shown therein, it consists of a solenoid assembly 51 also identified as 291 having a cylindrical core 294 which is provided with a centrally disposed bore 296. A plunger 297 is slidably mounted in the bore. A winding 298 is provided on the core 294. The solenoid assembly 291 is mounted on a bracket 301 which is secured to a plate

302 mounted within the control console. The plunger 297 is provided with a shaft 306 which is connected by a coupling 307 to a shaft 308 carried by the bellows assembly 291. The shaft 308 is a part of the bellows assembly 292. The bellows assembly 292 consists of a collapsible cylindrical member 309 in the form of a collapsible bellows of a suitable type such as those supplied by Bellofram. Means is provided for yieldably urging the piston 311 in a direction towards its home position and consists of a spring 312 carried within a housing 313 and having one end of the spring engaging the piston 311 to return the bellows to its normal or at home position. As shown in Figure 10, this tubing 314 is connected to a pressure transducer 316. It is also connected into the catheter 118 and also to the input of the solenoid operated valve V1 also identified as valve 317. The outlet from the valve 317 is connected to a tube 318 which is connected to two one-way check valves 319 and 321. The check valve 319 is open to ambient whereas the check valve 321 is connected through a filter 322. The filter 322 is connected to the outlet of a solenoid operated valve V2, also identified as valve 323. The outlet of the valve 323 is also connected to a linear plenum 324 which is open to ambient or to the atmosphere. The inlet to the valve 323 is connected to a one-stage regulator 326 mounted upon a tank or container 327 containing a suitable gas as, for example, carbon dioxide or helium. The valve V2 supplies gas to the linear plenum 324 which feeds substantially zero pressure gas to the bellows 309 through the filter 322 through the check valve 321 through the valve V1 through the tubing 314 through the spring and into the bellows 309.

A stroke of approximately 1/4 inch is utilized for the solenoid 51 to ensure there is adequate force applied by the solenoid 51 to the bellows 309. However, it should be appreciated that if desired a longer stroke can be utilized. When the solenoid assembly 291 is energized, the piston 311 of the bellows 309 is moved to create a positive pressure against the yieldable force of the spring 312 forcing gas into the catheter 118 and into the balloon 119. As soon as the solenoid assembly 291 is deenergized, the piston 311 is returned to its home position by the spring 312 which causes deflation of the balloon 119.

The solenoid operated valves V1 and V2 ensure that proper positive and negative pressures are provided within the bellows-balloon pneumatic circuit connected to the balloon 119. The solenoid valves V1 and V2 are pulsed with pulses having a width ranging from 10 to 20 milliseconds at the beginning of each inflation and deflation cycle for the balloon 119. The symbols for the valves V1 and V2 as shown in Figure 10 indicate that both of the

valves have two positions, in one position they are open, in the other position they are closed. For both valves when the 10 to 20 millisecond pulses are supplied they are set to be in the open position. When the pulses are not supplied, the valves are in the closed position.

The one-stage regulator 320 can provide a desired pressure between 1 and 50 psi. The linear plenum 324 is in the form of a long coiled tube which serves as a linear path for the gas so that whatever gas comes from the pressurized cylinder 327 will go through and exit to ambient to ensure that ambient air will not be drawn into the pneumatic circuit. The point at which the filter 322 meets the linear plenum 324 is a substantially zero pressure junction and is the point where gas from the cylinder 327 is transferred through the one-way valve 321 into the bellows balloon pneumatic circuit. The transfer of gas into the bellows-balloon pneumatic circuit is at substantially zero ambient pressure so that any pressure that is generated from the pressurized cylinder 327 carrying the gas cannot accidentally overpressurize the balloon 119 which is in the patient.

This transfer of gas can be understood by examining the timing diagram which is shown in Figure 11. As shown in Figure 11 from top to bottom there is shown an ECG trace 331, a pump stroke trace 332, a V1 and V2 trace 333, an S1 trace 334, and a balloon pressure trace 336. A typical balloon inflation deflation cycle can last for approximately 2/5 (two fifths) seconds which would be equivalent to approximately 150 beats per minute, as for example, as a maximum down to approximately less than 1 cycle per second or less than 60 beats per minute. This cycle time is completely dependent on the ECG as represented by the trace 331. As shown by traces 332 and 333 at the beginning of the pump stroke, the solenoid operated valves V1 and V2 are pulsed for a period of time ranging from 10 to 20 milliseconds. This time is adjusted to provide the desired peak pressure in the solenoid pneumatic circuit. After closing of the solenoid operated valves V1 and V2, the solenoid S1 which is the solenoid assembly 291 is energized to cause movement of the piston 311 to compress the gas within the bellows balloon pneumatic circuit to create a pressure equivalent to approximately 60 millimeters of mercury to cause inflation of the balloon 119 as shown by the trace 336 when inflation of the balloon is desired.

At the time that deflation of the balloon 119 is desired, for example, at the peak 337 of the R-wave trace 331, the valves V1 and V2 are energized for a predetermined period of time ranging from 10 to 20 milliseconds to permit a slight amount of positive pressure within the pressurized balloon bellows pneumatic circuit to escape to am-

bient as indicated by the portion 339 of the trace 336, thereafter immediately deenergizing the solenoid V1 and permitting the spring 312 to return the piston 311 to its home position. This generates a negative pressure within the balloon-bellows balloon pneumatic circuit to approximately -30 millimeters of mercury in comparison to ambient. This negative pressure is held until the next inflation of the balloon 119 is desired. As soon as the next inflation of the balloon 119 is desired, the valves V1 and V2 are again pulsed to permit some of this negative pressure to escape as indicated by the portion 341 of the balloon pressure trace 336.

As soon as some of this negative pressure has been relieved, the solenoid S1 is again energized to create a positive pressure. This sequence is repeated continuously with a pulsation of the valves V1 and V2 stabilizing the generation of the positive and negative pressures over time. During this sequence of operation, the bellows balloon pneumatic circuit is flushed with the desired carbon dioxide or helium gas. This is accomplished by pulsation of the valve V2. During each cycle the valve V2 is pulsed for a short time to fill the linear plenum 324 with the desired gas. As this linear plenum 324 is filled with a volume of this gas from the container 327 any ambient air within the plenum 324 is exposed to ambient through the exhaust 325. At this point in time there is essentially zero pressure at the junction between the linear plenum 324 and the filter 322. During the time when there is negative pressure within the balloon bellows pneumatic circuit which is at the time that the piston 311 is being returned to its home position by the spring 312, zero pressure gas is transferred from the plenum 324 through the check valve 321 through the bellows balloon pneumatic circuit. At the end of the pneumatic cycle when there is high pressure in the balloon bellows circuit created by operation of the solenoid S1, a small amount of this pressurized gas which is a mixture of the beginning gas and the carbon dioxide or helium gas in the plenum 324 which has been mixed with an exit through the valve V1 and through the check valve 319 to ambient air.

It should be understood that when the gas inflation apparatus is placed in operation, the apparatus will begin with ambient air and with each stroke, additional carbon dioxide or helium gas supplied by the container 327 is introduced. After a number of cycles, a sufficiently high concentration of the desired gas, as for example, carbon dioxide or helium is attained in the balloon pneumatic circuit to be acceptable medically. The longer the gas inflation apparatus operates, the higher the percentage of the carbon dioxide gas or helium achieved within the bellows pneumatic circuitry until it eventually approaches approximately 100%. To initially

facilitate flushing the ambient air with the desired gas, a temporary flush mode is provided by opening both V1 and V2 while cycling the solenoid S1 several times. This modality takes advantage of the full cycle time during which negative and positive pressures are generated in the bellows with respect to the exhaust and the gas input points for accelerating the exchange of resident air with the desired gas.

The stability of the balloon pressure over time is determined by the pulse width of the pulses on the valves V1 and V2. The valve V2 is pulsed at the same time as valve V1 because it is not critical when valve V2 is pulsed as long as some of the gas in the container 327 is transferred into the linear plenum 324. The timing of the operation of the valve V1 is important with the two cycles of operation of the valve V1 being labeled A and B with A being at the beginning of the stroke of the solenoid S1 and B being at the end of the stroke solenoid S1 which determines the long term stability of the pressures generated in the balloon. Thus if the length of the pulse time for the pulse A is increased, the pressure will increase in the balloon 119. Conversely if the time for the pulse A is decreased, the pressure in the balloon 119 will decrease. Similarly if the pulse time for the pulse B is increased the pressure in the balloon will decrease and if the width of pulse B is decreased, then the pressure in the balloon will increase. The reason for this is readily apparent. If a greater amount of negative pressure is exhausted then positive pressure, then over time the pressure will climb. Conversely, if a greater amount of positive pressure is exhausted than negative pressure, then over time the balloon pressure will fall. Thus it can be seen by adjusting the pulse width of the pulses A and B, the desired balloon pressure can be set and can be maintained over time.

The circuitry which is utilised for operating the solenoid operated valves V1 and V2 and the solenoid S1 is shown in Figure 12. The timing diagram for this circuit is shown in Figure 13. As shown in Figure 12, the circuitry consists of three one shot multi vibrators 351, 352 and 353 which can be of a conventional type as, for example, TTL circuitry such as the LS123 supplied by Texas Instruments. The output A of the multivibrator 351 is supplied to an OR gate 354 and also to the input of the multivibrator 353. The output 349 from the microprocessor 206 is supplied to the inputs of the two multivibrators 351 and 352. It is also supplied to an opto isolator 356 which has its output supplied to a field effect transistor 357 which has its output connected to the winding 298 of the solenoid S1. The output A of the multivibrator 351 is supplied to the input of the multivibrator 353 which has its output B supplied to another opto isolator 358 through a

field effect transistor 359 to the winding 298 of the solenoid S1. The field effect transistor 357 is provided with a five volt supply 361 through a diode 362. The field effect transistor 359 is provided with a 24 volt supply 363. The output C of the multivibrator 352 is supplied to the OR gate 354 which is connected to an opto isolator 364. The opto isolator 364 is connected to a field effect transistor 366 to the windings 367 and 368 of the solenoid operated valves V1 and V2. As can be seen the other ends of the windings 298, 367 and 368 are connected to ground. The field effect transistor 366 is also provided with a 24 volt supply 369.

The output 349 for the microprocessor is indicated in the timing diagram in Figure 13 as a curve 371. The curves at various points in the circuitry which are indicated as points A, B, C, D and E are shown as curves 372, 373, 374, 376 and 377 respectively.

The multivibrators 351, 352 and 353 are wired so that they are edge-triggered, that is, with either an up edge or a down edge providing a trigger to provide a pulse of the desired length. Thus when the microprocessor signal 349 supplies a pulse as shown by curve 371 and the edge of a pulse is ascertained by the multivibrator 351, its output A as indicated by the curve 372 provides a first pulse which is supplied through the OR gate 354 through the opto isolator 364 and the field effect transistor 366 to supply a voltage of the windings 367 and 368 to energize the valves V1 and V2. Thus it can be seen that the valves V1 and V2 are energized on the up edge of the pulse 381 from the microprocessor to provide a pulse 382 of desired length as, for example, from 10 to 20 microseconds for energizing the valves V1 and V2 as hereinbefore described.

At the same time as the up edge of the microprocessor pulse 381 occurs, a signal is supplied through the opto isolator FET to supply a five volt signal to the winding 298 of the solenoid S1. However, this voltage is not sufficient to actuate the solenoid S1 and therefore it will not be actuated with this voltage. This voltage is indicated by the step 383 in the curve 376.

The output from the multivibrator 351 is also supplied to the multivibrator 353 which is triggered on the negative going edge of the pulse 382 to provide the pulse 384 in the curve 373 from the output B of the multivibrator 353. This pulse 384 is of a suitable width as, for example, 50 milliseconds which is supplied through the opto isolator 358 and the FET 359 to supply 24 volts to the winding 298 of the solenoid S1 which will cause energization of the solenoid S1 to cause it to operate to advance the piston 311 in the manner hereinbefore described. This pulse 384 has a suitable width as, for example, 50 milliseconds. The 24 volts which is

supplied to the winding 298 of the solenoid S1 is represented by the step 386 in curve 376.

The third multivibrator 352 is triggered by the negative going edge of the pulse 381 of the microprocessor signal and its output C as shown by the curve 374 produces a pulse 387 having a suitable width as, for example, 10 to 20 milliseconds to supply a signal through the OR gate 354 through the opto oscillator 364, the FET 366 to the windings 367 and 368 of the valves V1 and V2. The output E from the FET 366 is shown by the curve 377 in which the pulses 388 correspond to the pulses 382 and the pulses 389 correspond to the pulses 387.

From the foregoing it can be seen that the solenoid S1 is operated to move the full amount of its travel during the 50 millisecond pulse 384. At the end of the 50 milliseconds, the voltage on the solenoid drops down to 5 volts which is sufficient to hold the solenoid in place for the duration of the cycle as indicated by the pulses 381 of the microprocessor.

In Figure 14, there is shown another embodiment of a pneumatic circuit which can be utilized for creating a balloon pressure without the use of a pressurized gas, as for example, from the container 327 as in connection with the embodiment previously described. As shown in Figure 14, this gas inflation apparatus utilizes the same bellows 309 with the piston 311 to supply air to an output tube 314 which is connected to the catheter 118 and the balloon 119. The pressure transducer 316 is also provided. The tubing 314 is connected to a solenoid operated valve V3 which is similar to the solenoid operated valve V1 or V2 as hereinbefore described with the exception that it is a three-way solenoid operated valve rather than a two-way solenoid operated valve. The valve V3 is connected to an extra plenum chamber 391 upon energization of the solenoid S1, a certain positive pressure of gas is generated of approximately 80 millimeters of mercury which pressure is created in the balloon 119 as well as in the extra plenum chamber 391 through the open valve V3. Upon completion of the upstroke of the piston 311, the solenoid V3 is closed to isolate the pressurized volume VE in the extra plenum 391. The pressurized gas in the extra plenum 391 is then permitted to escape to ambient through the valve V3 through the filtered exhaust 393. The volume VB in the bellows as well as in the balloon is at this point isolated from the extra plenum chamber 391. Upon the reverse stroke of the solenoid S1 and the movement of the piston 311, a negative pressure is created because the volume VE in the extra plenum 391 has been isolated from the bellows balloon pneumatic circuit. Once this negative pressure has been generated following the first upstroke, upon each downstroke

cycle the negative pressure will again be created because of the volume of gas which has been removed by the isolated extra plenum 391. Assuming there are no leaks in the system, operation of the apparatus can be sensed by the pressure transducer 316 and a signal is supplied to the microprocessor which supplies a signal to the solenoid operated valve V3 to connect the plenum 391 to the tubing 314 to remove gas from the bellows balloon pneumatic circuitry and then closing the solenoid operated valve V3 to isolate the extra plenum 391 from the bellows pneumatic circuitry at the end of the upstroke.

After this initialization, the appropriate positive-negative pressures can be maintained within desired limits. As explained previously, if this initialization is not maintained, the apparatus can be reinitialized. Thus by switching the extra plenum 391 into and out of the pneumatic circuitry, it is possible to generate the desired negative and positive pressures. By switching the extra plenum 391 into and out of the pneumatic circuitry, it is possible to generate a specific negative pressure from the specific stroke of the solenoid and the bellows-balloon pneumatic circuitry.

If it is desired to generate additional negative pressure the size of the extra plenum can be increased or alternatively, the apparatus can be cycled additional times, as for example, two or three times in succession to obtain the desired negative pressure. By cycling the extra plenum into and out of the pneumatic circuitry, it is possible to change the positive and negative destination pressures within the bellows balloon pneumatic circuitry. The balloon pressure generation apparatus shown in Figure 14 makes it possible to inflate and deflate the balloon 119 when filtered air is adequate. The pneumatic circuitry permits the generation of the desired positive and negative pressures without the use of a vacuum pump and without relying upon the use of pressurized gasses.

It has been found that the pneumatic systems utilized for inflating and deflating the balloons in a positive and negative manner is advantageous over the use of an auto-inflatable balloon hereinbefore described. It does not require a specific volume of blood to be used each time of balloon inflation. By way of example, utilizing the auto-inflatable balloon and assuming the balloon has a volume of one-half of a cubic centimeter on an 8 or 10 millimeter balloon at 120 beats per minute uses 60 cubic centimeters of blood during each minute for inflating and deflating the balloon. This volume of blood travels to and from the disposable pump and in the balloon catheter during each cycle instead of going out the tip and being utilized as flow for retroperfusion. Utilizing a gas as the inflation medium for the balloon, this volume of blood which was pre-

viously used for balloon inflation and deflation can now be passed out of the tip of the catheter and increase the flow for retroperfusion. Alternatively, the pumping pressure or the cycling of the pump can be decreased.

It also has been found that the auto-inflatable balloon is sometimes undesirable because it is possible for the pressure of the blood in the coronary sinus to backfill the balloon through the tip valve and to cause undesirable high pressures in the coronary sinus. With the gas inflation system for the balloon, this cannot occur because the balloon is independently controlled and inflated and deflated. Also it is held deflated in a positive manner so that there cannot be an accidental buildup of coronary sinus pressure.

An additional safety factor over an auto-inflatable balloon is provided by the fact that the coronary sinus pressure does not increase beyond the balloon inflation pressure of approximately 60 millimeters of mercury. Should conditions exist to generate a greater pressure, the balloon will automatically decrease in size to allow this greater coronary sinus pressure to decrease and equilibrate with the balloon pressure even without active deflation of the balloon with the microprocessor.

The circuitry which is shown in Figure 8 for the previous embodiment has been slightly revised as shown in the circuitry shown in Figure 15. The keyboard 171 has been eliminated as has the heart rate display 181 and the flow display 182 which have been replaced by the touch panel 268 and the flat panel display 266.

An integral blood flow meter has been incorporated into the console 12 which can be of a conventional type and is provided with a probe 401 which is connected to patient isolation circuitry 402 to a flow amplifier processing and probe excitation circuitry 403 to provide a flow signal to the microprocessor 206.

In addition, a pressure transducer output from a coronary sinus pressure transducer 406 (see Figure 1) of a conventional type is supplied to the cable 199 on the console 12. It can be identified as "coronary sinus pressure in". The transducer 406 is connected by tubing 407 to a y fitting 408 connected to a three lumen catheter 118a. The cable 199 is connected to an amplifier 411 which supplies an analog sinus pressure indication to the analog digital converter 207 to the microprocessor controller 206. This makes it possible to utilize sinus pressure to automatically control the blood pumping and/or balloon inflation period. The output from the amplifier 411 is also supplied to another amplifier 412 to an output terminal 413 to a cable 414 which can be identified as "coronary sinus out".

A patient ECG apparatus 201 has been incor-

porated into the console 12 so that a direct isolation amplified connection can be made to the patient through the patient electrodes 203. The cable 202 in Figure 8 is used for making direct contact to electrodes 203 placed on the patient and can be identified as "ECG in". The auxiliary ECG input 191 has been retained in the event it is desired to utilize an external ECG monitor and is identified as "AUX ECG input".

As also can be seen the gas inflation mechanism 271 has its input connected to the output of the microprocessor 206 for the purposes hereinbefore described.

The cable 197 is adapted to be connected to the auxiliary ECG and can be identified as "AUX ECG in". Conduit 198 is connected to the pump and can be identified as "Pump Stroke". Conduit 198 is connected to the tubing 314. The tubing 314 is also connected to a fitting 416 mounted on a side wall of the console 12. An extension tube 417 connects to the fitting 416 to another y fitting 418 connected to the three lumen catheter 118a.

The operation of the system and apparatus for retroperfusion is readily apparent from the description hereinbefore given with respect to the previous embodiment. The use of the apparatus for retroinfusion may now be briefly described as follows. Let it be assumed that a typical drug such as streptokinase is to be delivered systemically and it is desirable to reduce the amount of the drug delivered to the patient. By delivering the same directly to the heart via the coronary sinus it is possible to achieve the same positive coronary effect while utilizing less volume of the drug and therefore decreasing the systemic side effects. This can be readily accomplished with the present apparatus by introducing the drug into the container 276 and having the same delivered directly to the coronary sinus during each pump cycle. By way of example, the balloon can be inflated and held inflated during the time that the diluted drug is being pumped continuously or nearly continuously into the coronary sinus until the coronary sinus reaches a certain limit at which time the balloon can be deflated and the pumping of the drug terminated.

Still another embodiment of the retroperfusion and retroinfusion apparatus and system is shown in Figures 16-20 that is particularly adapted for use in cerebral retroperfusion and which also can be utilized for coronary retroperfusion wherein balloon inflation occurs independently of the pumping of the blood for improving the flow pressure characteristics of the pumped blood.

In the apparatus and system shown in Figures 16-20, a bladder type pump cassette 426 is utilized in place of the piston type cassette utilized in the previous embodiments hereinbefore described. The pump console 12 which is shown in Figures 16 and

17 is in many respects similar to the pump consoles 12 described in connection with the previous embodiments. The bladder-type pump cassette 426 is mounted on the side of the console 12 rather than on top as with the previous piston-type pump cassettes. The bladder-type pump cassette 426 consists of a housing 427 formed of a suitable material such as a transparent plastic to provide a cylindrical portion 427a to provide a cylindrical bladder chamber 428. The housing also consists of another smaller inverted bottle-like generally cylindrical type portion 427b which defines a blood level chamber 429. A web 430 is provided for interconnecting the two portions 427a and 427b.

A bladder 431 which has a generally cylindrical configuration is disposed within the chamber 428. The bladder 431 is formed of a suitable flexible material such as plastic or rubber. The lower extremity of the bladder 431 is mounted on a flanged inlet fitting 432 which has a flow passage 433 extending therethrough. The flow passage 433 is adapted to be occluded by a ball valve 434 of a suitable material such as stainless steel. The ball valve 434 normally is seated within a conical valve seat 436 which is formed in an insert 437 of a relatively soft material such as silicone rubber. The ball valve 434 is movable from a closed position in engagement with the valve seat 436 to an open and uppermost position where it engages a bar 438 extending the passage 433.

The insert 437 is carried by an end cap 439 which is mounted in a flanged end portion 441 of the cylindrical portion 427a by suitable means such as ultrasonic welding. The upper extremity of the bladder 431 is thickened as shown particularly in Figure 9 and is provided with a radially extending flanged portion 431a. The portion 431a is retained between a flanged end portion 442 provided on the cylindrical portion 427a of the housing 427 and by a flanged outlet fitting 443. The fitting 443 is held in place by a retaining ring 444 and is bonded to the fitting 443 and the flanged end portion 442 by ultrasonic welding. The upper thickened portion of the bladder 431 forms a conical valve seat 446 which accommodates a ball valve 447 formed of a suitable material such as stainless steel. The ball valve 447 is movable between a closed position which occludes a passage 448 provided in the bladder 431 and an open position in which the ball valve 447 is away from the valve seat 446 with its uppermost travel being limited by a bar 449.

An outlet tube 451 is mounted within the outlet fitting 443 and is retained therein by suitable means such as shrink tubing 452. An inlet tube 453 is mounted in the inlet fitting 439 and is in communication with a flow passage 454 provided in the insert 437. The inlet tubing 453 which also can be described as a transfer tube and is connected in

the lower extremity of the portion 427b and is in communication with the bubble chamber 429. An inlet tube 456 is also connected to the bottom extremity of the portion 427b. The inlet tube 456 is also in communication with the blood level chamber 429.

An inlet port 459 is provided on the cylindrical portion 427a and is in communication with the portion of the chamber 428 between the cylindrical bladder 431 and the wall of the portion 427a. The inlet port 459 is provided for introducing a relatively incompressible liquid into the space between the exterior wall of the bladder and the interior wall of the cylindrical portion 427a. A removable cap 461 covers the inlet 459. A cylindrical extension 462 is provided on the portion 427a and extends generally at right angles thereto. The cylindrical extension has mounted therein a piston 463. The piston 463 is provided with an annular recess 464 which has a piston seal 466 disposed therein. A piston rod 467 is formed integral with the piston 463 and is provided with a head 468 which is adapted to be secured to the pump actuator of the controller 12 as hereinafter described. An injection site plug 457 is mounted in the upper extremity of the portion 427b of the housing 427 and is formed of a suitable material such as rubber which can be penetrated by a needle. The plug 457 closes the upper extremity of the cylindrical portion 427b.

The cassette 426 is mounted on the controller 12 in a suitable manner such as by a clamping assembly 471. The clamping assembly 471 is comprised of a base plate 472 which can be secured to the control console 12 in a suitable manner such as by screws (not shown). Upper and lower clamping members 473 and 474 are provided which are adapted to engage the pump cassette 426. The lower clamping member 474 is secured to the base plate 472 in a fixed position and extends outwardly at right angles therefrom. The top surface of the upper clamping member 473 is engaged by a knob 476. A threaded rod 477 is mounted in the knob 476 and extends through a hole 478 provided in the upper clamping member 473. A C-ring 479 is mounted on the rod 477 and engages the lower surface of the upper clamping member 473 to permit rotation of the threaded rod 477 within the hole 478 but limiting vertical movement of the rod 477 relative to the upper clamping member 473. The lower extremity of the rod 477 is threaded into a threaded hole 480 provided in a support plate 481 which is secured to and extends perpendicularly from the base plate 472. The clamping members 473 and 474 are provided with slots 484 extending inwardly toward the control console from the outer extremities of the same which are adapted to receive the upper and lower extremities of the pump cassette 426. They are

also provided with circular recesses 486 to receive the upper and lower extremities of the pump cassette.

Another U-shaped member 491 is mounted on the side wall 129 adjacent the base plate 472 and extends perpendicularly from the side wall 29. The U-shaped member is provided with a U-shaped recess 492 which is adapted to receive the cylindrical portion 427b of the housing 427.

The rack and pinion arrangement herein before described and shown in Figure 2 is utilized in the controller 12 as shown in Figure 17 with the principal difference being that the rack and pinion assembly is rotated by 90° so that it can operate the side mounted bladder type pump cassette 426, rather than the piston type pump cassette 81 hereinbefore described.

Cassette position means 158a similar to the cassette positioning means hereinbefore described is provided for ascertaining when the pump cassette 421 is properly positioned. The pump console 12 consists of a microswitch 159a which is mounted within the pump console and which is adapted to be operated by an operating arm or rod 160a which extends through the U-shaped member 491 and terminates in the U of the U-shaped recess 492 of the U-shaped member 491 and is adapted to sense when the portion 427b is disposed within the recess 492.

The rack and pinion arrangement hereinbefore described and is shown in Figure 2 is utilized in the controller 12 as shown in Figure 17 with the principal difference being that the rack and pinion assembly is rotated by 90° so that it can operate the side mounted bladder-type cassette 426 rather than the piston-type pump cassette 81 hereinbefore described. As shown in Figure 17, the rack 61 extends through hole a 494 and is coupled to the head 468 of the piston 463.

In utilizing the bladder-type pump cassette 426, a relatively incompressible liquid is utilized for filling the space in the chamber 428 which is not filled by the bladder 431. Typically this space can be filled with a sterile saline solution 496 by inserting it through the inlet 459 and then closing the same with the cap 461.

The pump cassette 426 can then be filled with blood in a conventional manner. The pump cassette 426 can then be mounted in the clamp assembly 471 by raising the upper member 473 to receive the pump cassette so that it can be positioned within the circular recess 486. The knob 476 can then be rotated to bring the upper member 473 downwardly to securely clamp the pump cassette 426 between the upper and lower members 473 and 474. When the pump cassette is positioned in this manner, the head 468 is positioned within the recess 111 of the rack 61. Similarly, the cylindrical

portion 427b is positioned within the U-shaped recess 492 of the U-shaped member 491 of the operating rod 160a to operate the micro-switch 159 to indicate that the pump cassette 426 is properly positioned with respect to the controller 12. As the pinion 63 is driven, it can be seen that the rack 61 is reciprocated inwardly and outwardly with respect to the side wall 129 to cause operation of the piston 463. Operation of the piston 463 causes compression of the liquid saline solution 496 which in turn causes compression of the flexible bladder 431 which causes blood to flow outwardly to the flow passage 448 by raising the ball 447 to cause blood to flow through the outlet tubing 451 to the patient. On the return stroke, the ball valve 447 moves to a closed position and the ball valve 434 is opened to withdraw blood from the chamber 429 through the transfer tube 453. Additional blood is brought in to the chamber 428 by the inlet tube 456. The blood level within the chamber 429 is sensed by the blood level detector 145 which is provided in the U-shaped member 491 in the same manner it is provided in the members 143 in the embodiment shown in Figure 1. Similarly, a pinch-off valve mechanism 151 of the type hereinbefore described is provided for pinching off the flow of blood in the tube 456 when that is necessary.

It can be seen that in the embodiment of the bladder-type pump cassette shown in Figures 18 and 20 that an intermediate sterile saline driving or liquid solution 496 is interposed between the piston 463 and the isolating silicone bladder 431 which carries blood. It is desirable that the driving solution 496 utilized in the pump cassette be a solution which in the event a leak should develop in the bladder 431 and mix with the blood would be harmless to the patient. This is one of the reasons why a sterile saline solution is utilized for the intermediate captive driving solution.

The inlet chamber 429, in addition to serving as a chamber for blood level detection also serves to deaerate and aspirate a trapped gas in the blood in the same manner as in the embodiments previously described.

In operation of the pump console 12 when the piston 463 is driven in a forward direction positive pressure is created within the cylindrical portion 427a and in the bladder 431 and causes the ball check valve 434 at the base of the cassette 426 to be moved to a closed position and to cause opening of the ball check valve 447 at the top of the cassette 426 to drive blood out of the bladder 431 through the tubing 451. When the piston 463 is driven in a rearward or reverse position (toward the control console 12) a negative pressure is generated within the chamber 428 and in the bladder 431 to cause the ball valve 447 at the top of the

cassette 426 to be closed and to cause opening of the valve 434 at the base of the cassette 426 to bring blood into the bladder 431 from the reservoir 429 and to fill the bladder 431 in preparation for the next forward stroke of the piston 463.

The apparatus of the present embodiment may be utilized for performing retroperfusion in the same manner as in the previous embodiment of the invention hereinbefore described. However, it has been found desirable in certain applications to cause independent inflation of the balloon 119.

The outlet tube 451 is connected through the tubing clamp 117 by tubing 499 to the center arm of a three-arm adapter 501 which is connected by tubing 502 to a two-arm adapter 503. One arm of the adapter 503 is connected to a three lumen tubing 118a which is connected to the balloon 119. One side arm of the three-arm adapter 501 is connected to the tubing 407 which is connected to the transducer 406. The tubing 417 is connected to a two-arm adapter 504. One arm of the two-arm adapter 504 is connected by tubing 506 to the other side arm of the three arm adapter 501.

It can be seen that with the apparatus shown in Figure 16 that retroperfusion in the manner hereinbefore described can be carried out for use in the coronary vessels of the heart. In connection with such retroperfusion it has been found that it often is desirable to maintain inflation of the balloon 119 for more than one cardiac cycle. This can be readily accomplished by the controller 12 since the balloon is gas inflated independently of the cyclical pumping of the blood. It has been found that this creates increased pressure in the coronary sinus which causes the retroperfusion of arterial blood to penetrate more deeply into the ischemic regions.

In addition to the apparatus and system being useful in connection with retroperfusion and retroinfusion of the coronary vessels by occluding the coronary sinus and pumping blood into the coronary veins in synchronization with the electrocardiogram it is also possible to occlude one or both of the cerebral sinuses and to pump blood into the cerebral veins in a pulsatile or nearly continuous manner without synchronization with the electrocardiogram. The purpose is the same as with coronary retroperfusion, that is, to provide oxygenated blood and to wash blood in an ischemic region of the brain which has been rendered ischemic by the event of a stroke and by the subsequent occlusion of one or more of the cerebral arteries.

In connection with cerebral retroperfusion it has been found it is desirable to provide a second balloon 119a as shown in Figure 16. The human brain 511 as shown in Figure 16 is provided with cerebral venous system 512 which includes the Sigmoid sinus, Transverse sinus, Inferior sagittal sinus, the Superior sagittal sinus and branches

thereof. As shown such a venous system is provided with two major drainage paths from the Superior sagittal sinus 516 in the form of two jugular veins 513 and 514. In order to accomplish retroperfusion it is necessary to occlude the drainage paths from the brain. Typically, this can be accomplished by occluding both of the jugular veins 513 and 514 when retroperfusion flow is introduced into the coronary sinus through one or both of the jugular veins 513 and 514. This is particularly true if the retroperfusion flow is to reach beyond the bifurcation 517 occurring at the entrance to the Superior sagittal sinus 516. It is only necessary to introduce blood flow through one of the balloons as, for example, balloon 119 but inflation of both of the balloons 119 and 119a by the use of independent gas inflation is necessary in order to provide the desired occlusion of both of the jugular veins 513 and 514. It should be appreciated that if desired the drainage paths from the brain can be occluded higher up in the brain in other portions of the brain above the jugular veins. In cerebral retroperfusion, the balloons 119 and 119a are connected respectively by tubes 521 and 522 to adapters 503 and 504 respectively as shown in Figure 16. Both of the tubes 521 and 522 carry balloon inflation lumens. The balloon 119a is provided solely for the purpose of occluding the jugular vein 514. However, the balloon 119, in addition to providing occlusion for the jugular vein 513 also contains the infusion lumen for delivering arterialized blood as well as a pressure monitoring lumen communicating with the pressure transducer 406.

As explained previously, retroperfusion can be carried out by balloons independently inflated from the pumping of the blood for the purpose of attaining a better flow-pressure characteristic in the coronary or cerebral sinuses. For coronary retroperfusion, as hereinbefore explained, pumping is synchronized to the electrocardiogram and pumping occurs in the diastolic phase. However, inflation and deflation of the gas driven balloon 119 is controlled independently, either as hereinbefore described or inflated and deflated for several cardiac cycles or inflated and deflated for a predetermined time as, for example, several seconds. The length of time or number of cardiac cycles for the inflation and deflation is adjustable so that desired coronary sinus pressure is achieved with a particular selected flow rate. While the balloon 119 is deflated, the blood pumping may also be temporarily halted to allow for sinus drainage and reduction of coronary sinus pressure. For cerebral retroperfusion the pumping may or may not be synchronized to the electrocardiogram and the flow is less pulsatile than with the coronary retroperfusion. In cerebral retroperfusion the inflation and deflation of the gas driven balloons 119 and 119a is also controlled

independently from the activation of blood flow with the inflation and deflation being adjustable in length of time so that the desired cerebral sinus pressure is achieved with a selected blood flow rate.

It is apparent from the foregoing that there has been provided a retroperfusion and retroinfusion system, apparatus and method which has many advantages. The use of the gas inflation mechanism makes it possible to positively inflate and deflate the balloon within very precise limits. It is possible to introduce drugs into the coronary sinus with great efficacy eliminating the necessity of introducing excess amounts of the drug to obtain the desired effect without associated systemic side effects. The retroperfusion and retroinfusion system, apparatus and method in addition to being utilized in connection with occluding the coronary sinus and pumping blood into the coronary veins can also be utilized for cerebral retroperfusion and retroinfusion.

### Claims

1. In a control apparatus for supplying arterial blood of a patient to the venous side of the patient's heart, a pump having an inlet adapted to be coupled to an artery of the patient and an outlet adapted to be connected to a vein of the patient, movable means forming a part of the pump movable through a pump stroke for moving a liquid from the inlet to the outlet of the pump, a stepper motor, means coupling the stepper motor to the piston means of the pump for causing operation of the piston means, electronic circuitry for driving the stepper motor, the electronic circuitry including means for sensing the presence of and R wave in a patient and for operating the stepper motor in response to the sensed R wave.

2. Apparatus as in Claim 1 together with a bubble chamber coupled to the pump for receiving arterial blood and wherein said electronic circuitry includes detecting means associated with the bubble chamber and connected to the microprocessor for informing the microprocessor when bubbles of air appear in the arterial blood supply and also when the amount of arterial blood drops below a predetermined level to stop operation of the pump.

3. Apparatus as in Claim 1 wherein the microprocessor has been programmed to start the pump stroke at approximately 45% of the R wave to R wave interval and terminates at approximately 95% of the R wave to R wave interval and wherein the signals to the stepper motor are supplied during the upstroke at a considerably faster rate so as to provide a pause at the upper extremity of the pump stroke and prior to the downstroke so as to

permit any pressure buildup in the pump to dissipate and to permit a continued flow of blood to occur prior to the downstroke.

4. In a method for supplying arterial blood into a venous region of the heart of a patient, taking an electrocardiogram of the patient, ascertaining when an R wave is present in the electrocardiogram, supplying arterial blood from the patient under positive pressure into a venous region of the heart for a predetermined period of time in accordance with the R wave of the patient, providing a pause period for a predetermined period of time to permit any positive pressure created to be dissipated and to permit continued flow to occur and thereafter allowing antegrade flow of venous blood from the heart for a predetermined period of time in accordance with the R wave of the patient.

5. In a control apparatus for supplying arterial blood of a patient to the venous side of a patient's heart by the use of a catheter having a distal extremity, an inflatable balloon carried by the distal extremity and having a balloon inflation lumen in communication with the balloon, a blood pump having an inlet adapted to be connected to a blood vessel of the patient and having an outlet adapted to be connected to another blood vessel of the patient for pumping the blood of the patient, motorized means for operating the blood pump and gas pressure generation means adapted to be connected to the lumen in communication with the balloon of the catheter and synchronized with the operation of the blood pump for inflating and deflating the balloon with a gas.

6. In a method for supplying arterial blood into a venous region of the heart by the use of a catheter having a distal extremity, an inflatable balloon carried by the distal extremity, a balloon inflation lumen in communication with the balloon and having a blood flow lumen therein through which blood can flow, supplying arterial blood from the patient under positive pressure through the blood flow lumen of the catheter at predetermined periods of time into a venous region of the heart and periodically inflating the balloon with a gas in synchronism with the supplying of arterial blood of the patient into the catheter.

7. In a control apparatus for supplying arterial blood of a patient to the venous side of a patient's organ, a balloon catheter having a distal extremity and an inflatable balloon carried by the distal extremity, the catheter having a balloon inflation lumen in communication with the balloon and a blood flow lumen extending through the balloon, a blood pump having an inlet adapted to be connected to an arterial blood vessel of the patient and having an outlet adapted to be connected to a venous blood vessel of the patient, motorized

means for operating the blood pump and gas pressure generation means adapted to be connected to the balloon inflation lumen.

8. Apparatus as in Claim 7 wherein the patient's organ is the patient's brain together with a second catheter having a distal extremity and an inflatable balloon carried by the distal extremity, the second catheter having a balloon inflation lumen in communication with the balloon, and wherein the gas pressure generation means is adapted to be connected to the balloon inflation lumen in communication with the balloon of said second catheter.

9. In a method for supplying arterial blood into a venous region of an organ of a patient by the use of a first catheter having a distal extremity and an inflatable balloon carried by the distal extremity, the catheter having a balloon inflation lumen in communication with the balloon and a blood flow lumen extending through the balloon, the method comprising placing the balloon in a venous vessel of the patient's organ, supplying arterial blood from the patient under positive pressure through the blood flow lumen of the catheter at predetermined periods of time into a venous region of the patient's organ and inflating the balloon with a gas.

10. A method as in Claim 9 wherein the organ is the brain and wherein the method is accomplished by the use of a second catheter having a distal extremity and inflatable balloon carried by the distal extremity, the second catheter having a balloon inflation lumen in communication with the balloon of the second catheter, the method comprising the additional steps of placing the balloon of the second catheter in another venous vessel of the patient's organ, synchronously inflating and deflating the balloons of the first and second catheters for occluding of venous flow simultaneously in two blood vessels of the patient's organ.

11. In a pump cassette for a liquid, a housing providing a chamber, a flexible bladder disposed in the chamber, inlet and outlet valve means carried by the housing and providing unidirectional flow of a liquid through the bladder, a substantially incompressible driving liquid disposed in the chamber and surrounding the bladder, and a movable piston carried by the housing for causing compression and decompression of the liquid in the chamber to cause liquid to flow through the bladder.

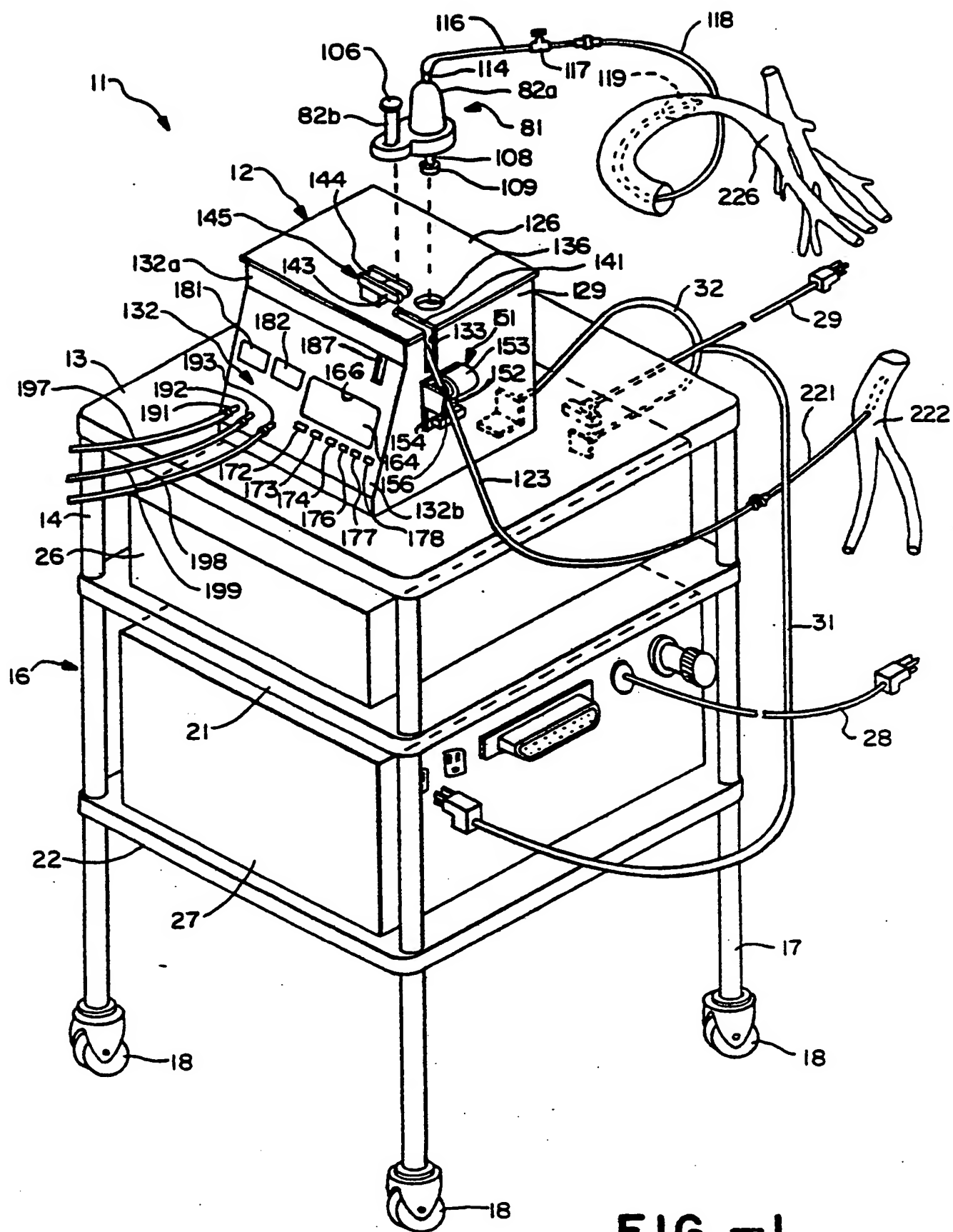
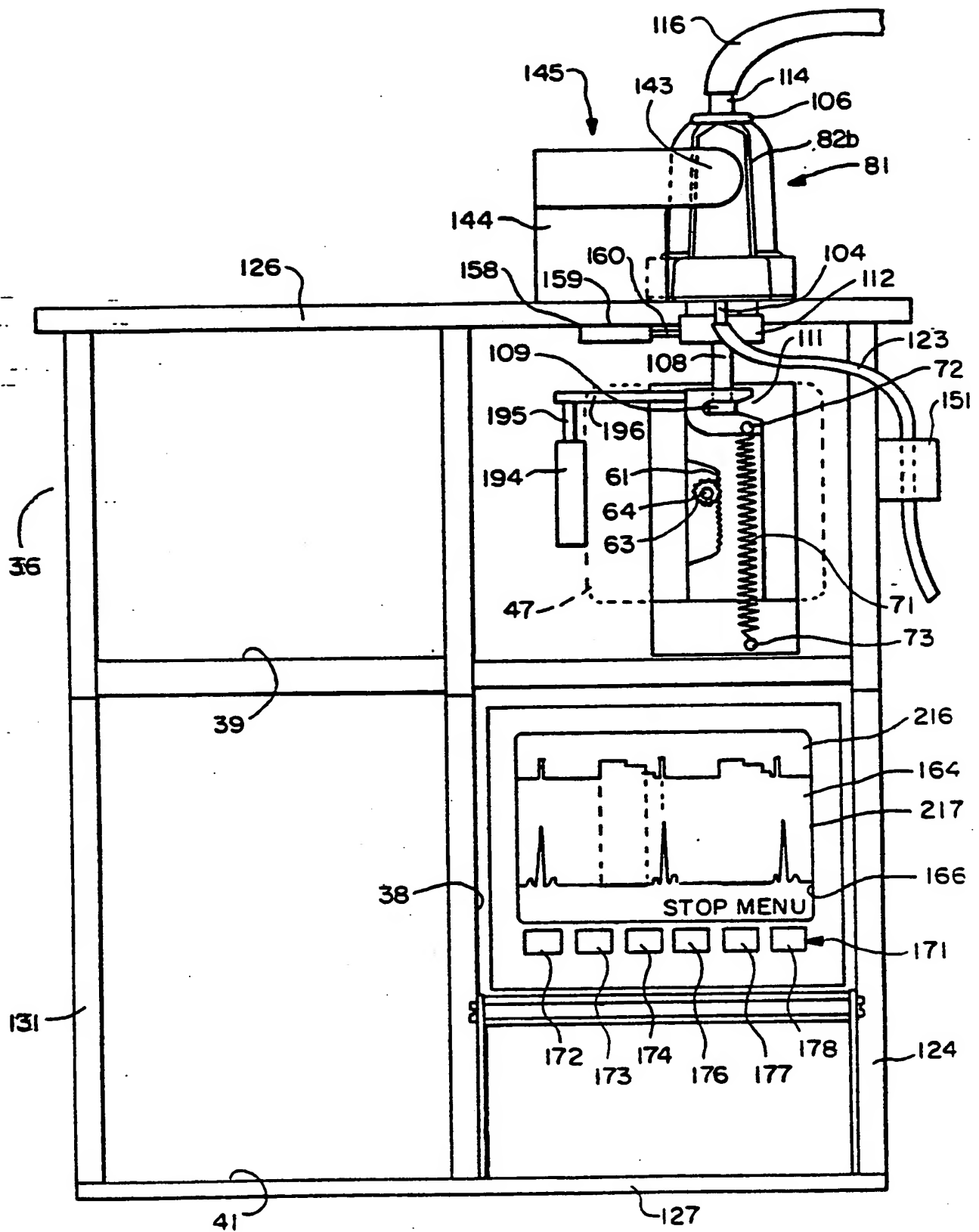


FIG. -1



**FIG. - 2**

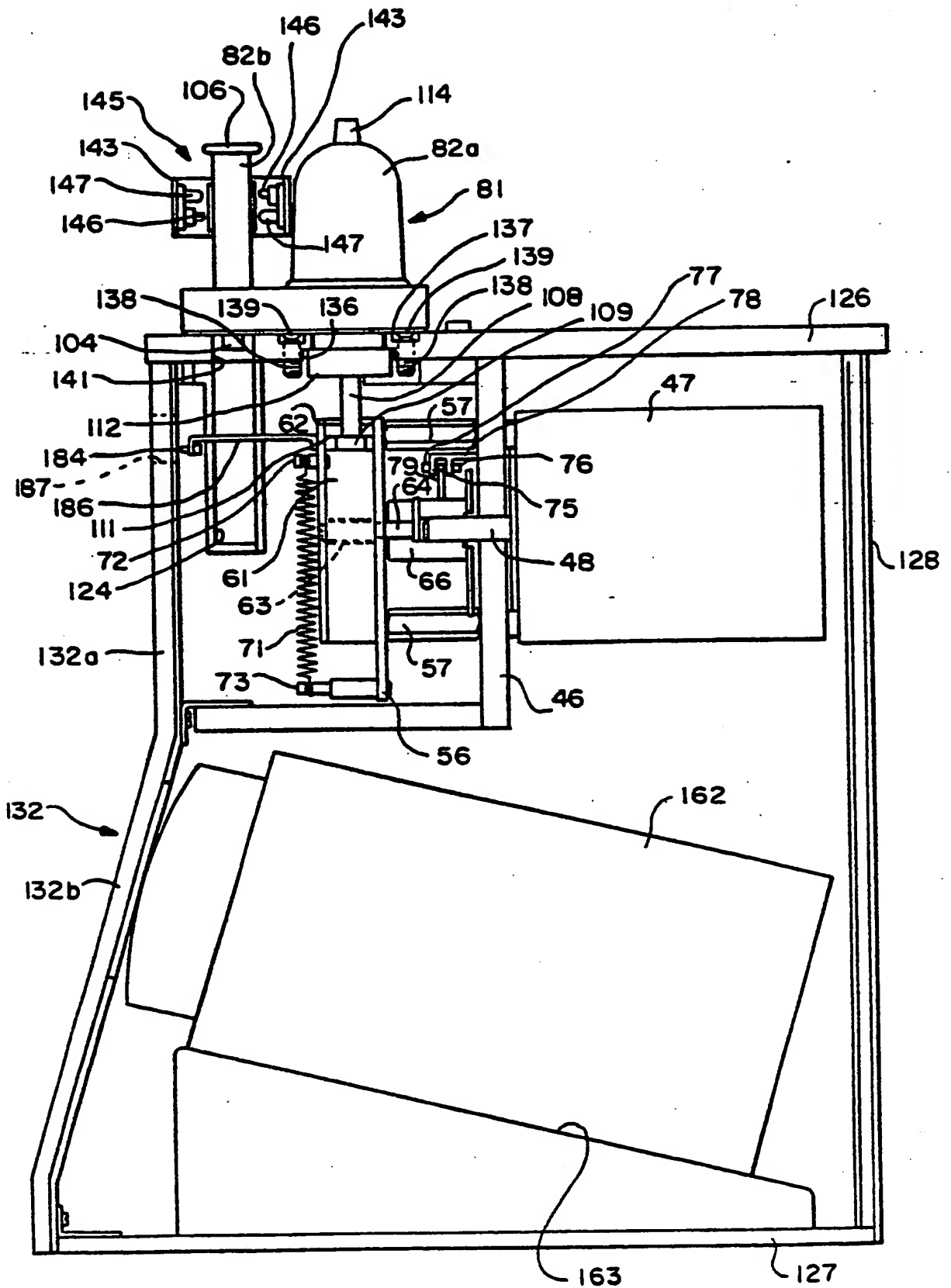


FIG. - 3

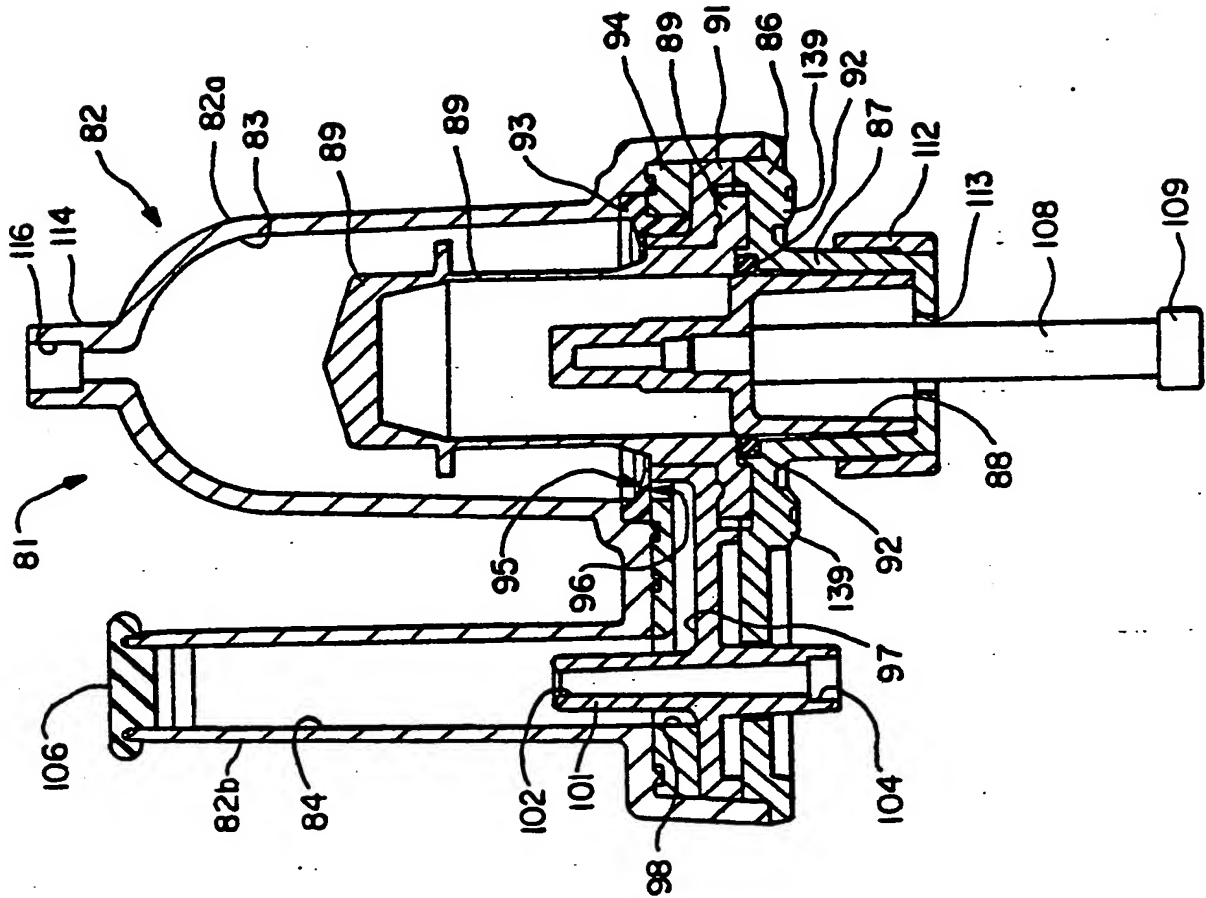


FIG.-4

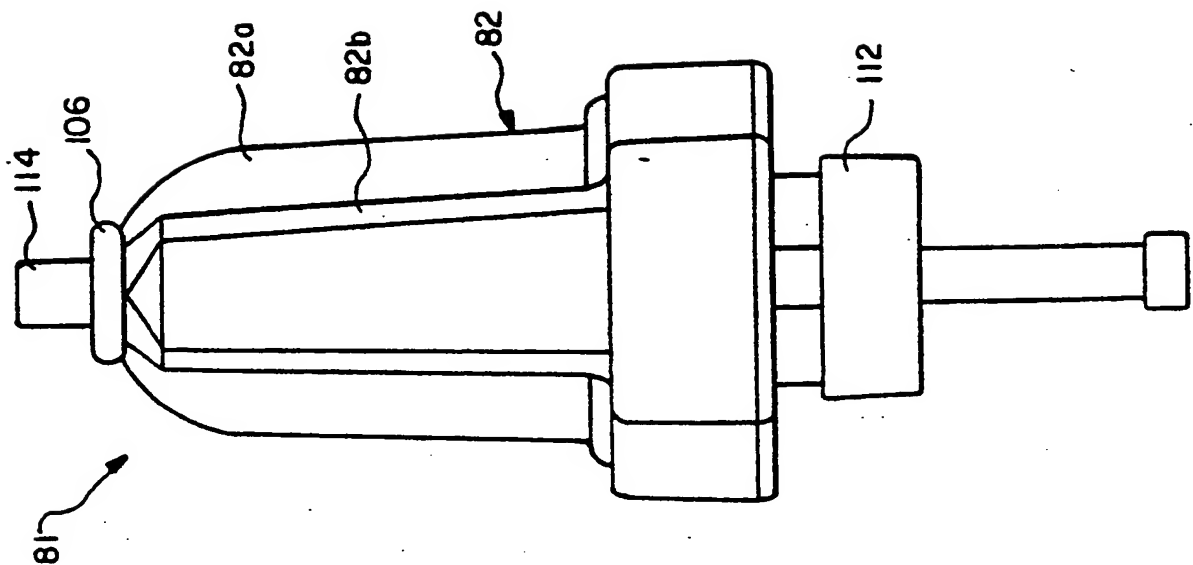


FIG.-5

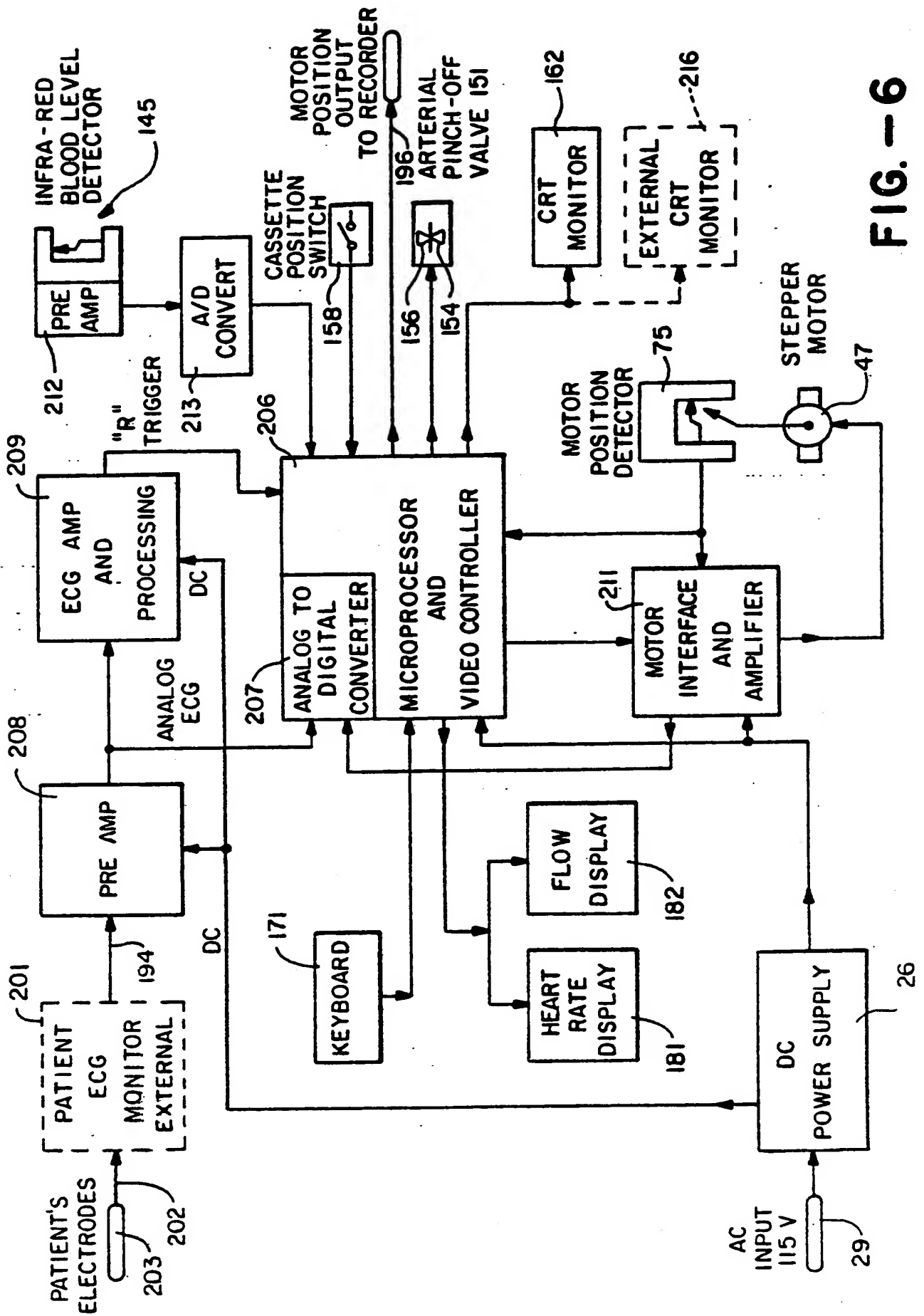


FIG. 6

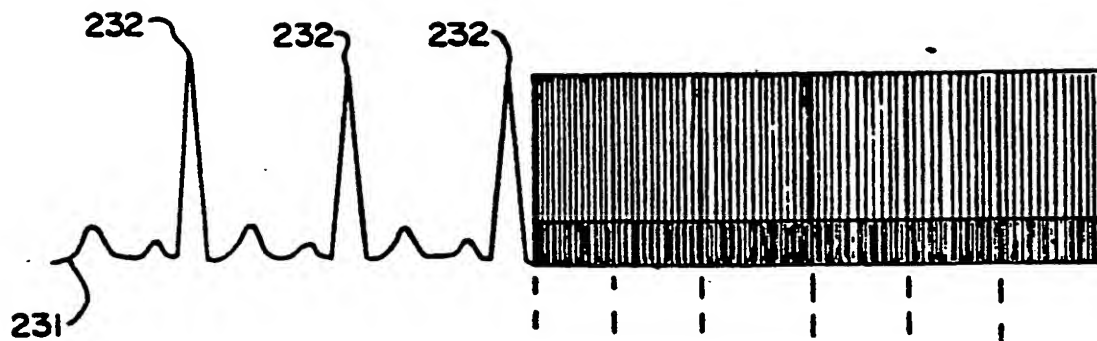


FIG. -7A

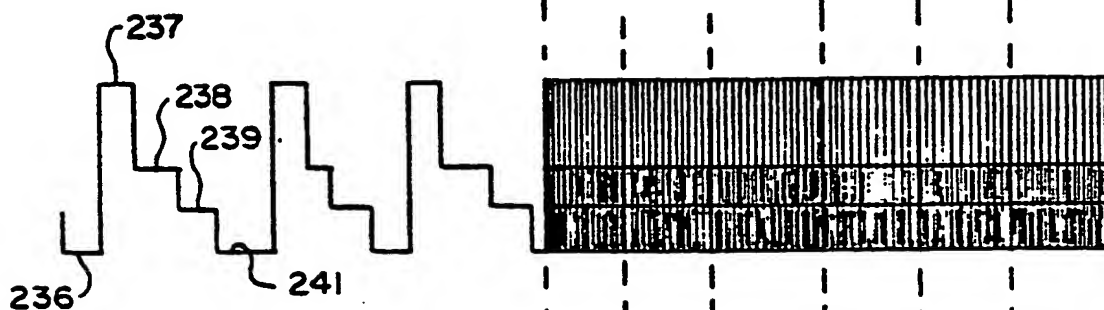


FIG. -7B

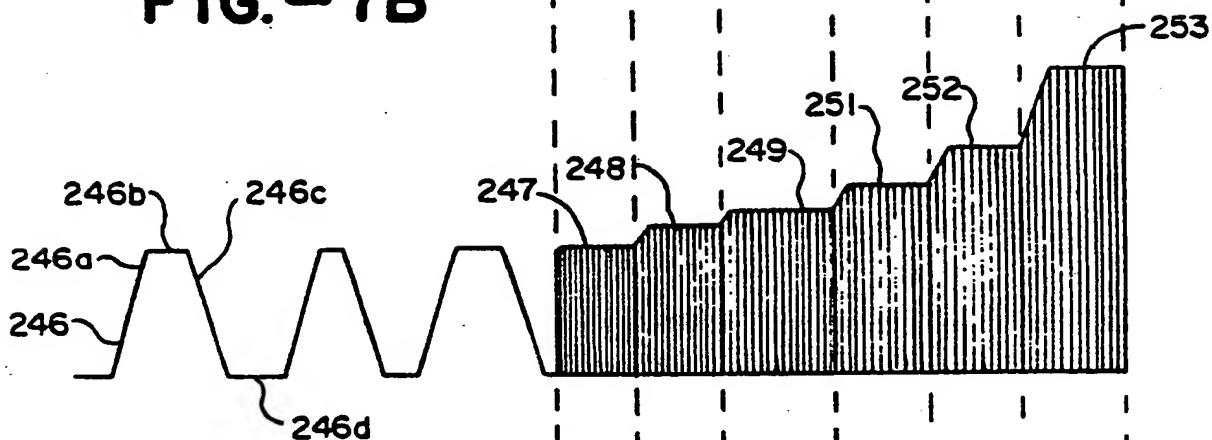


FIG. -7C

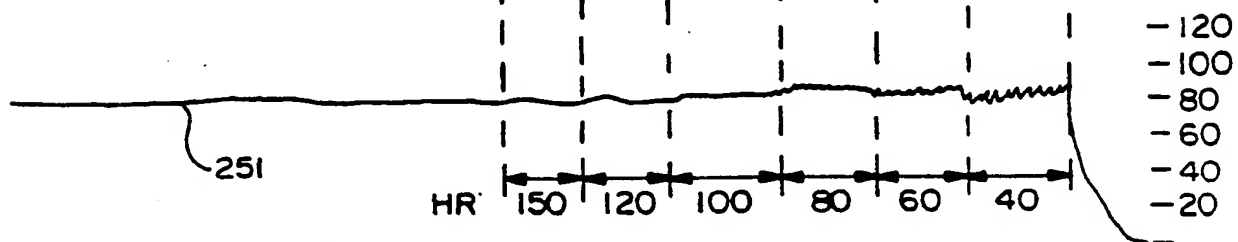


FIG. -7D

(FLOW SET = 80)

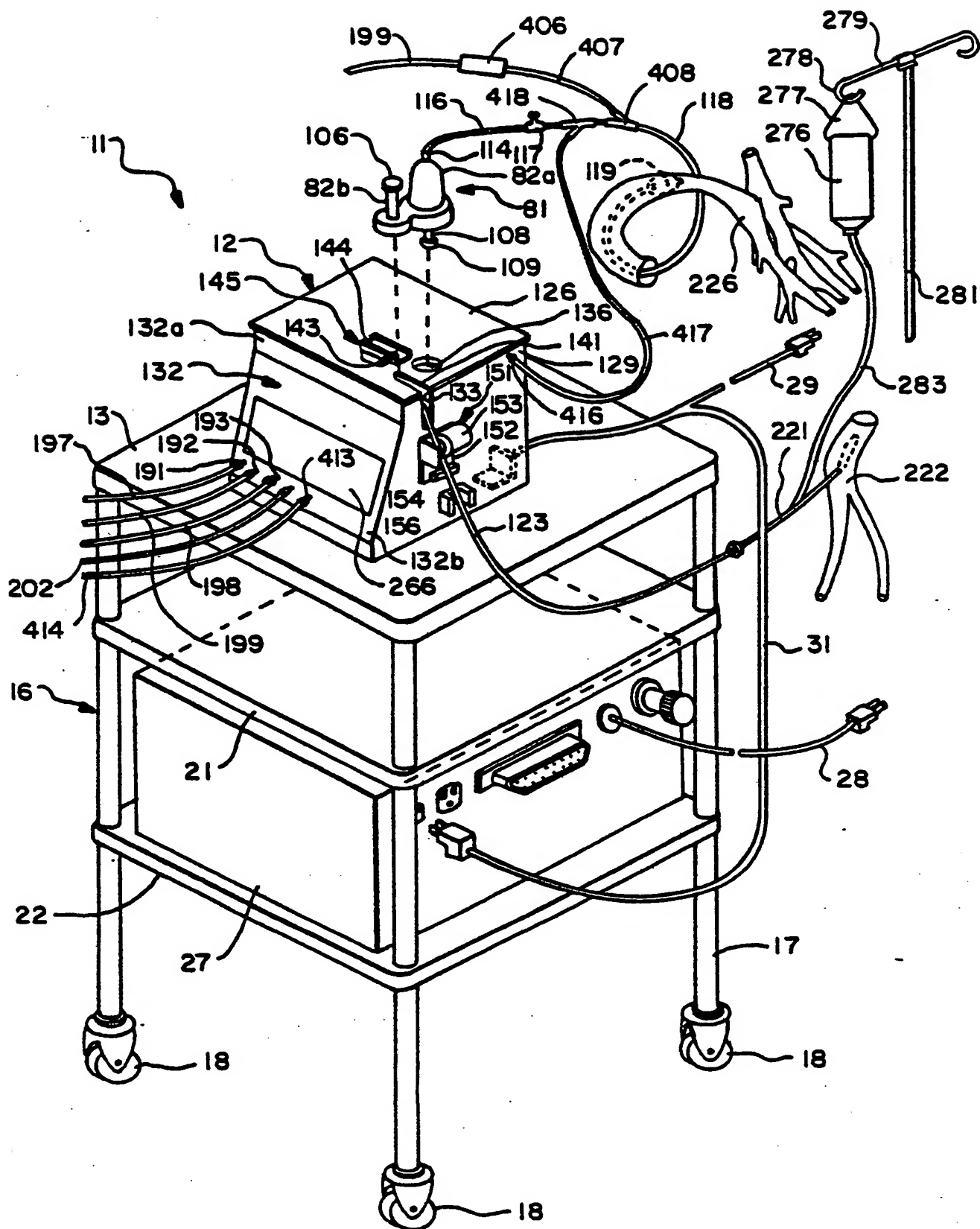


FIG. - 8

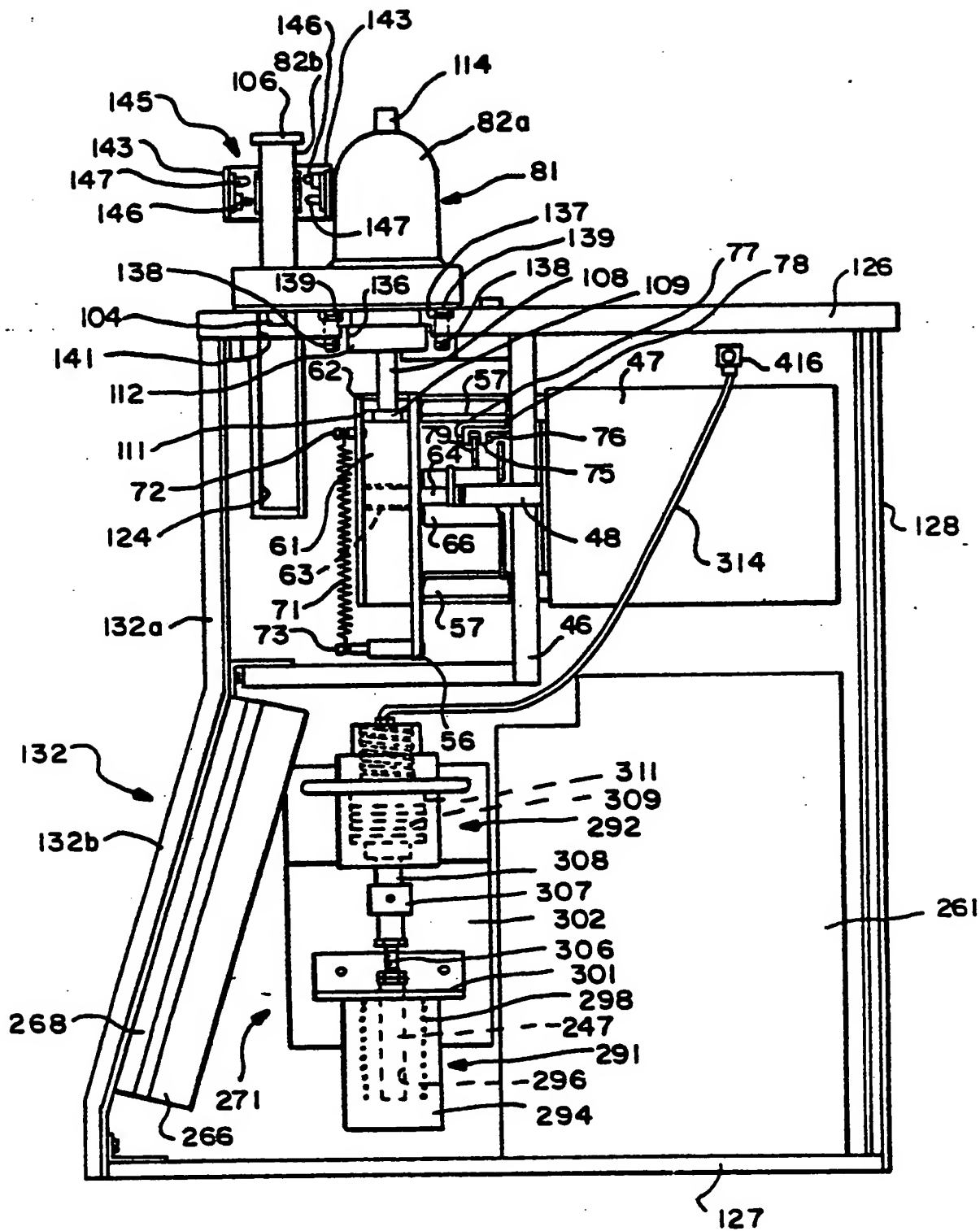


FIG.-9

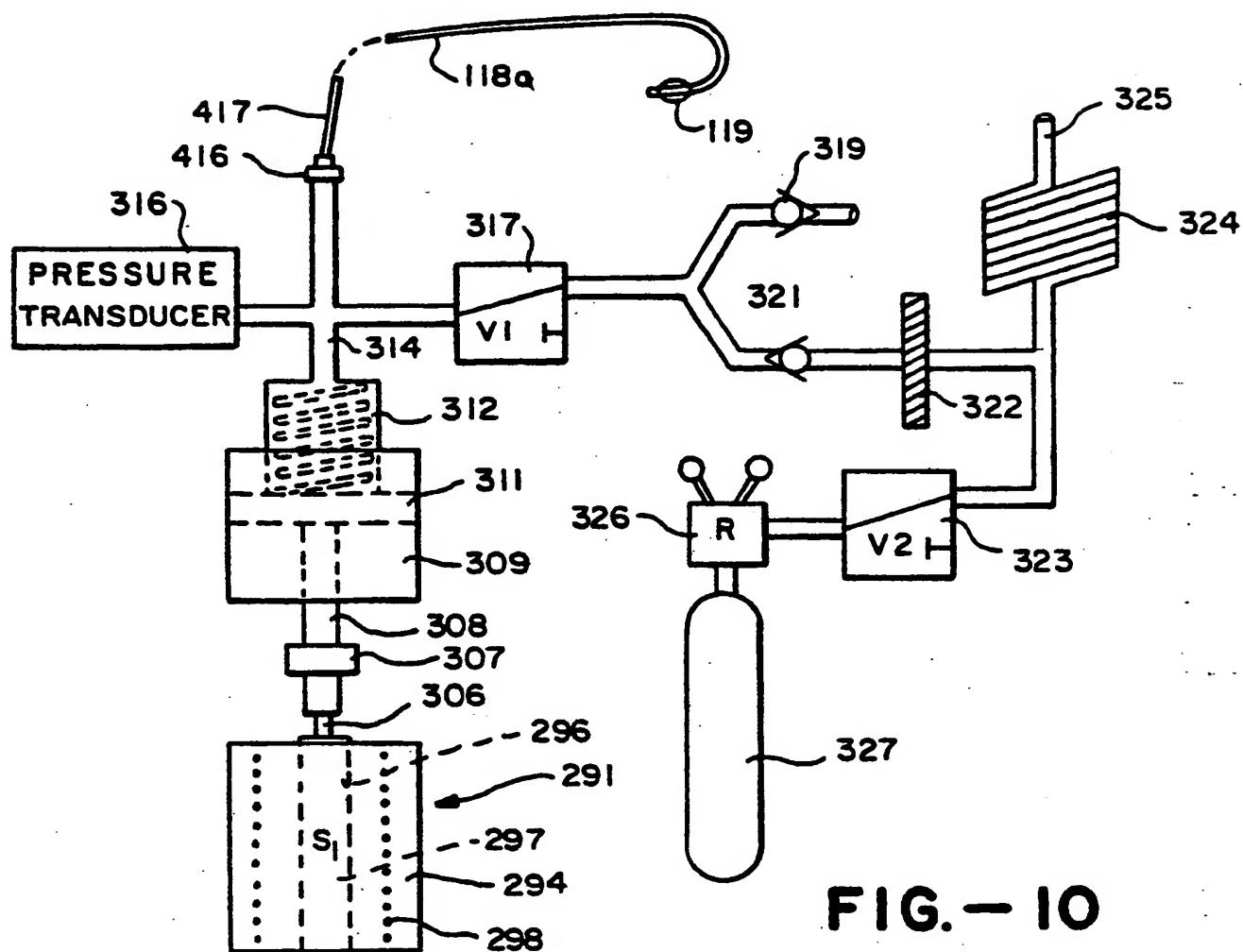


FIG. - 10

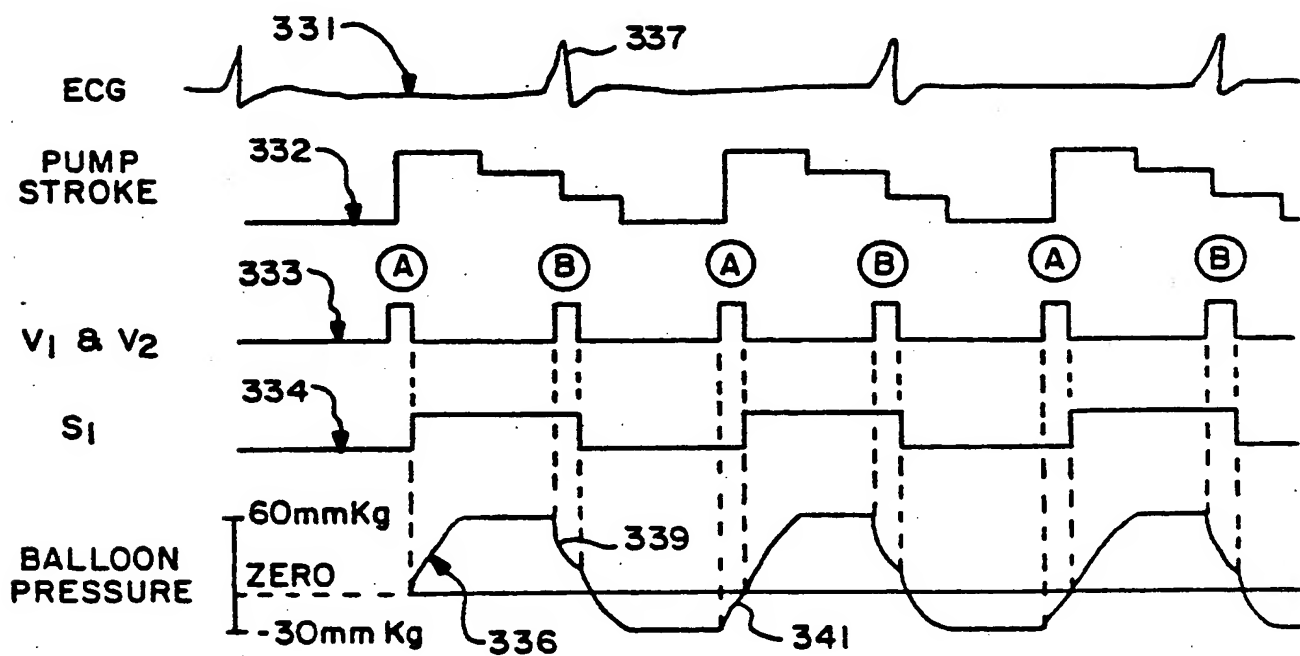
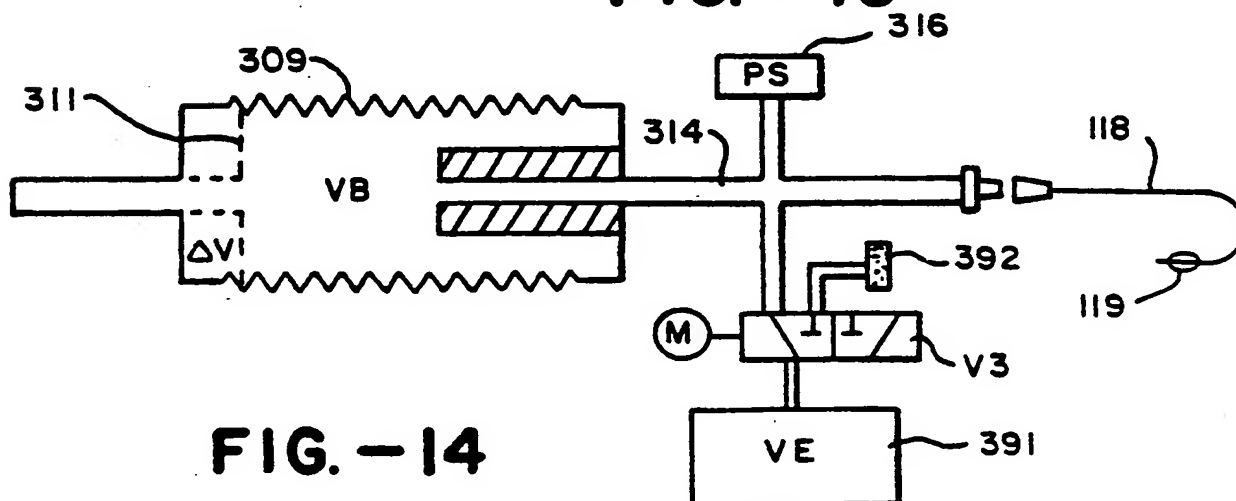
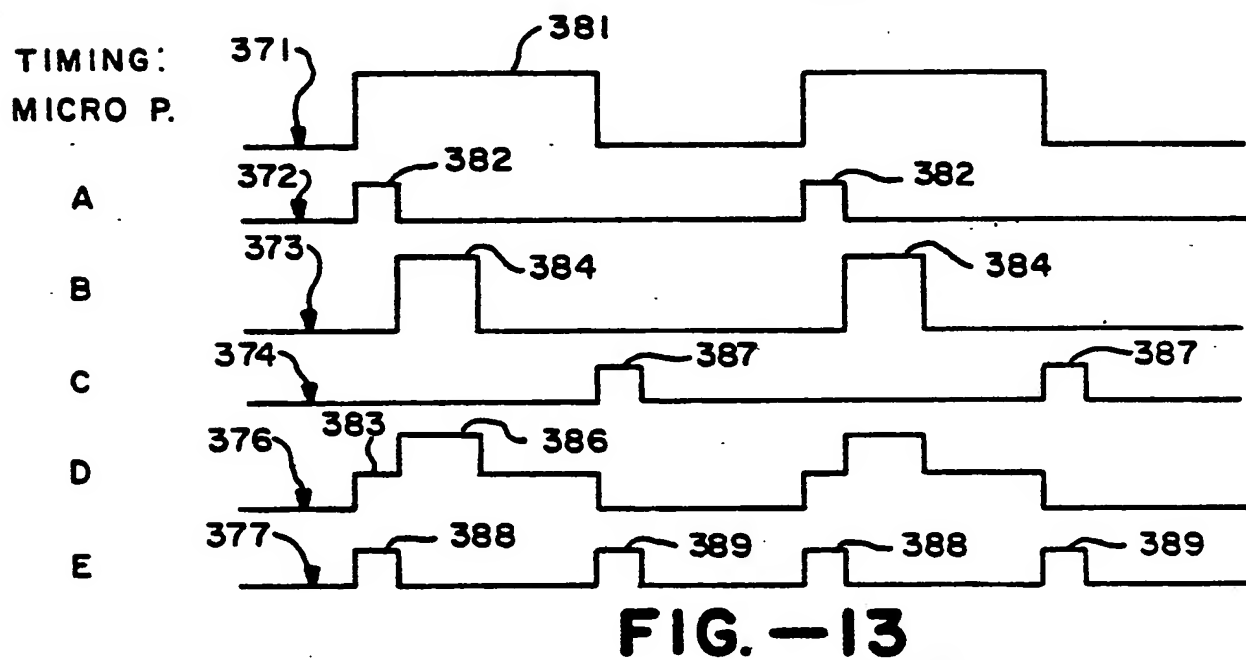
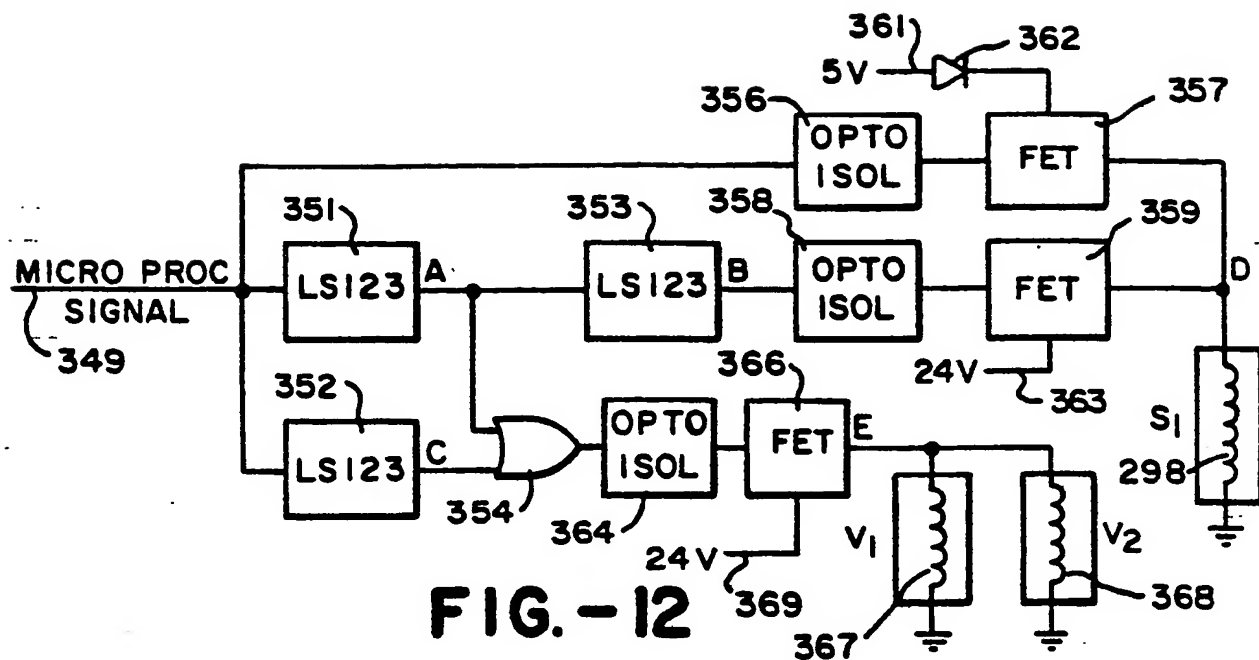


FIG. - 11



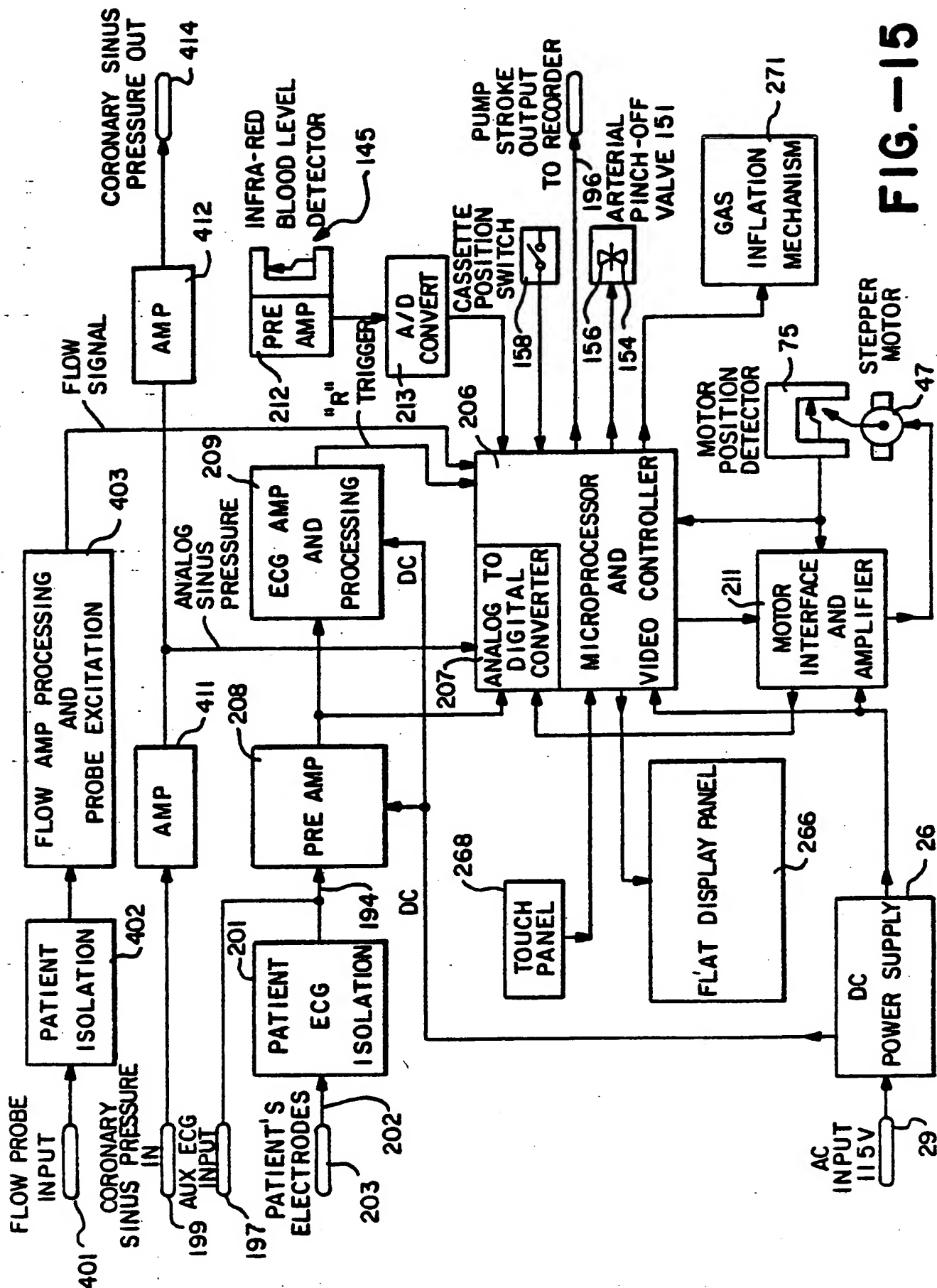


FIG. 15

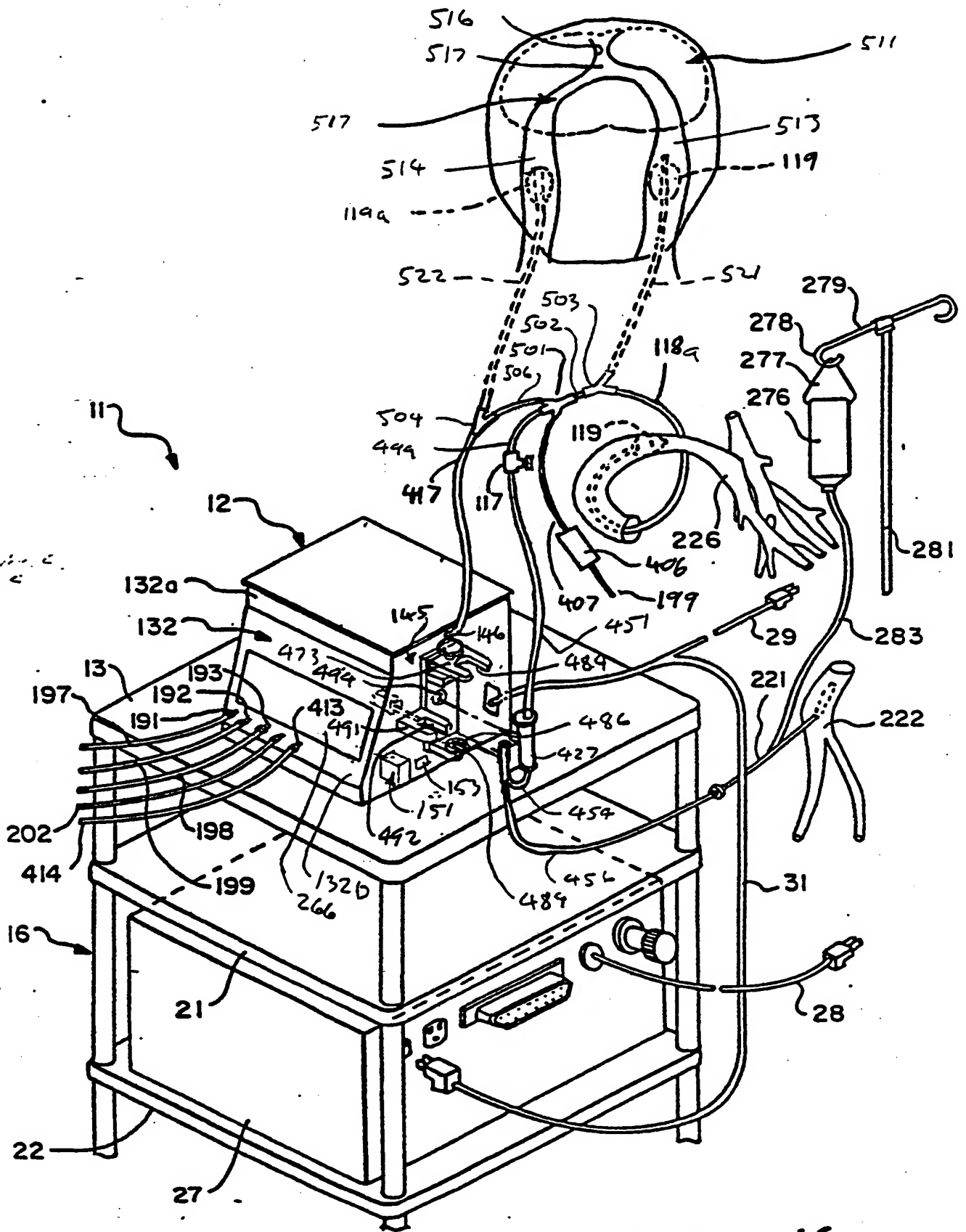


FIG. — 16

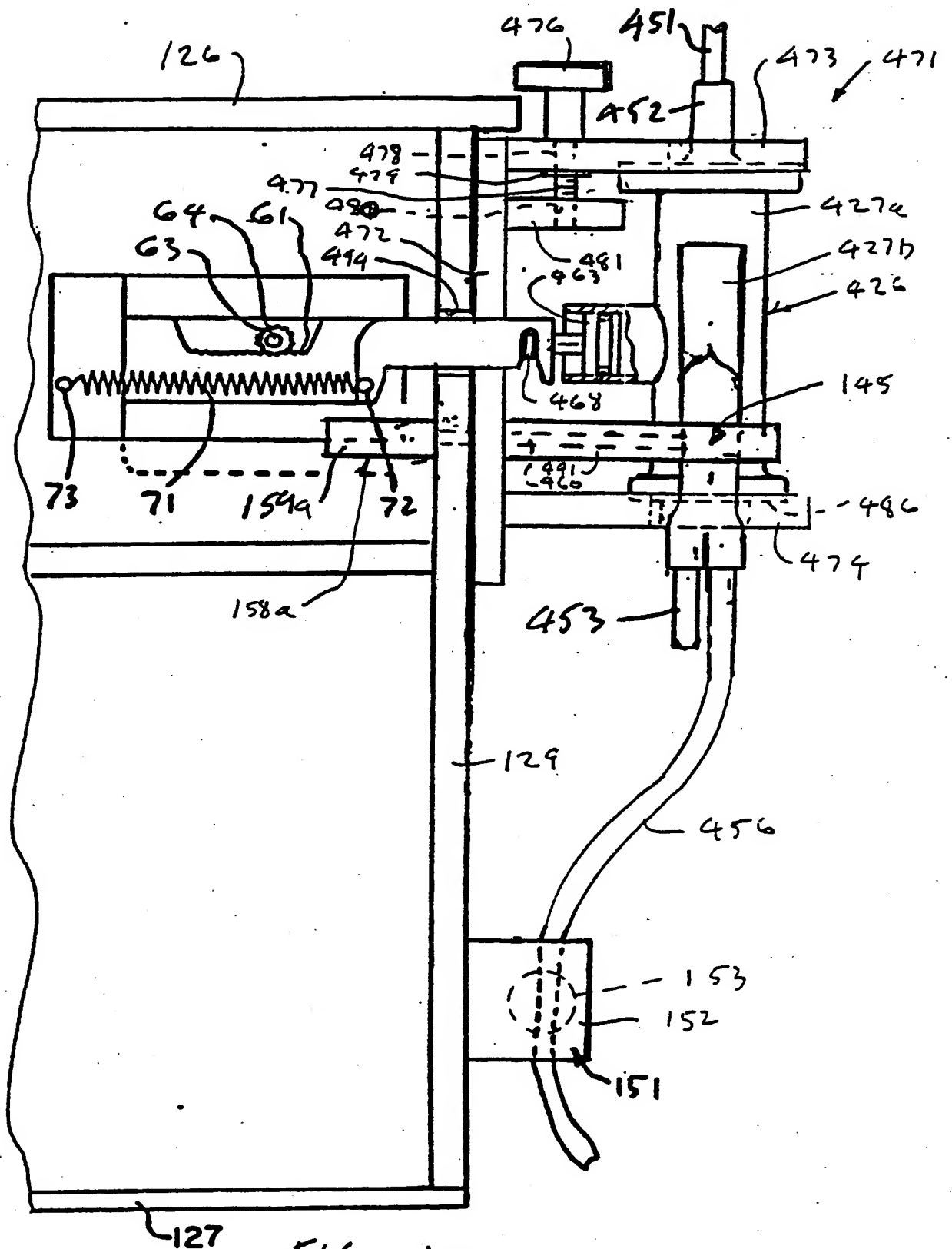


FIG-17

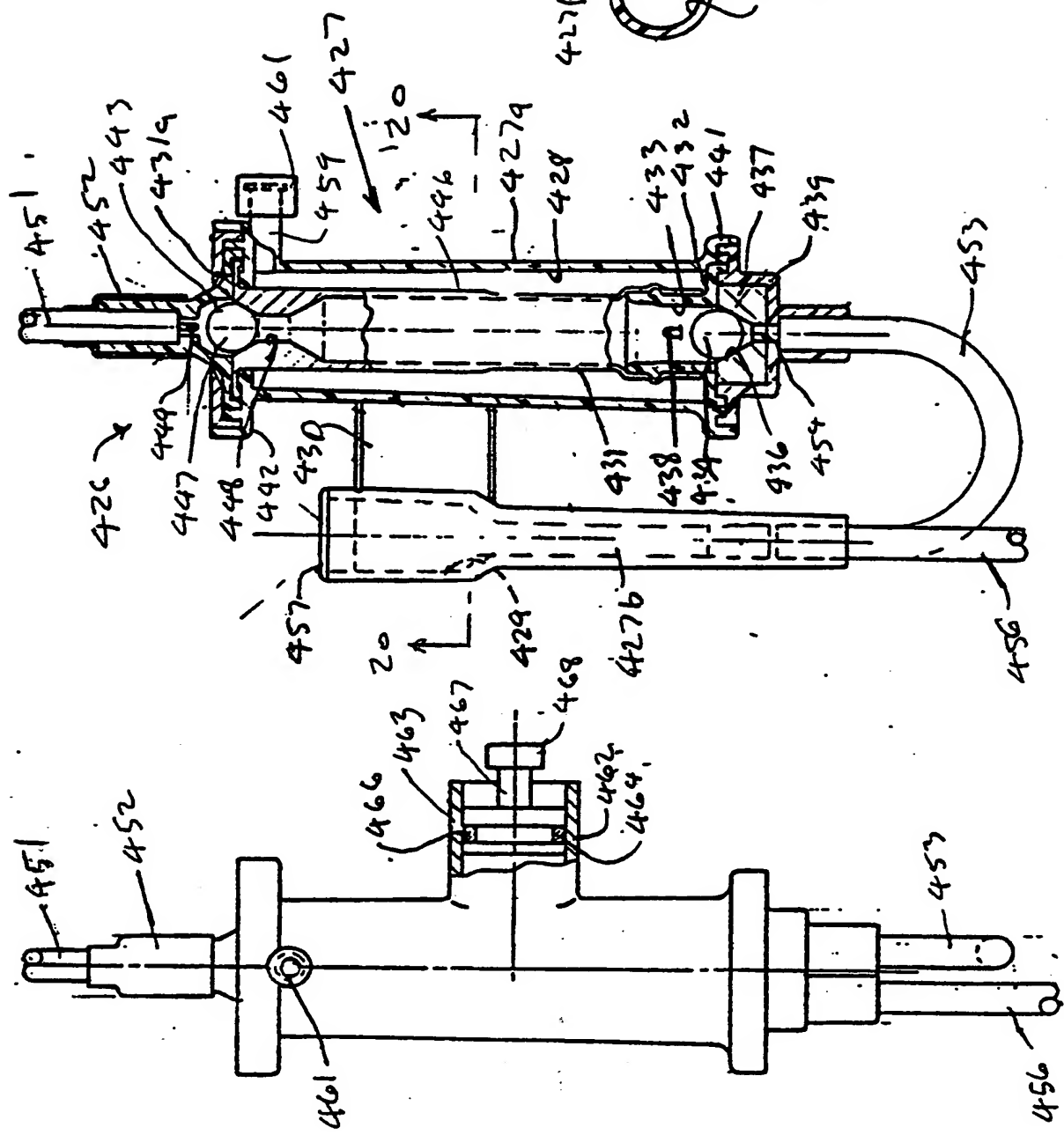


FIG. -19

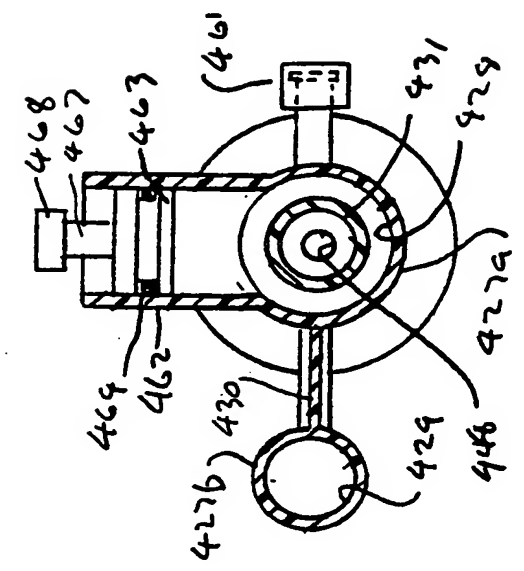


FIG. -20